

METHOD ARTICLE

Differential analyses for RNA-seq: transcript-level estimates improve gene-level inferences [version 1; peer review: 2 approved]

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Abstract

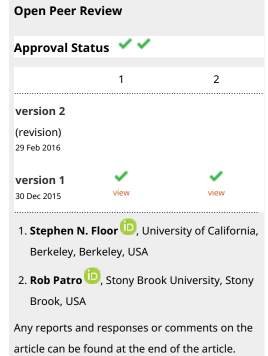
High-throughput sequencing of cDNA (RNA-seq) is used extensively to characterize the transcriptome of cells. Many transcriptomic studies aim at comparing either abundance levels or the transcriptome composition between given conditions, and as a first step, the sequencing reads must be used as the basis for abundance quantification of transcriptomic features of interest, such as genes or transcripts. Several different quantification approaches have been proposed, ranging from simple counting of reads that overlap given genomic regions to more complex estimation of underlying transcript abundances. In this paper, we show that gene-level abundance estimates and statistical inference offer advantages over transcriptlevel analyses, in terms of performance and interpretability. We also illustrate that while the presence of differential isoform usage can lead to inflated false discovery rates in differential expression analyses on simple count matrices and transcript-level abundance estimates improve the performance in simulated data, the difference is relatively minor in several real data sets. Finally, we provide an R package (*tximport*) to help users integrate transcript-level abundance estimates from common quantification pipelines into count-based statistical inference engines.

Keywords

RNA-seq, quantification, gene expression, transcriptomics



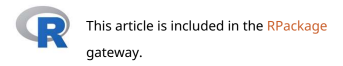
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Introduction

Quantification and comparison of isoform- or gene-level expression based on high throughput sequencing reads from cDNA (RNAseq) is arguably among the most common tasks in modern computational molecular biology. Currently, one of the most common approaches is to define a set of non-overlapping targets (typically, genes) and use the number of reads overlapping a target as a measure of its abundance, or expression level. Several software packages have been developed for performing such "simple" counting (e.g., featureCounts¹ and HTSeq-count²). More recently, the field has seen a surge in methods aimed at quantifying the abundances of individual transcripts (e.g., Cufflinks³, RSEM⁴, BitSeq⁵, kallisto⁶ and Salmon⁷). These methods provide higher resolution than simple counting, and by circumventing the computationally costly read alignment step, some are considerably faster. However, isoform quantification is more complex than the simple counting, due to the high degree of overlap among transcripts. Currently, there is no consensus regarding the optimal resolution or method for quantification and downstream analysis of transcriptomic output.

Another point of debate is the unit in which abundance is given. The traditional R/FPKM^{8,9} (reads/fragments per kilobase per million reads) has been largely superseded by the TPM¹⁰ (transcripts per million), since the latter is more consistent across libraries. Regardless, both of these units attempt to "correct for" sequencing depth and feature length and thus do not reflect the influence of these on quantification uncertainty. In order to account for these aspects, most statistical tools for analysis of RNA-seq data operate instead on the *count* scale. While these tools were designed to be applied to simple read counts, the degree to which their performance is affected by using fractional estimated counts resulting from portioning reads aligning to multiple transcripts is still an open question. The fact that the most common sequencing protocols provide reads that are much shorter than the average transcript length implies that the observed read counts depend on the transcript's length as well as abundance; thus, simple counts are arguably less accurate measures than TPMs of the true abundance of RNA molecules from given genes. The use of gene counts as input to statistical tools typically assumes that the length of the expressed part of a gene does not change across samples and thus length can therefore be ignored for differential analysis.

In the analysis of transcriptomic data, as for any other application, it is of utmost importance that the question of interest is precisely defined before a computational approach is selected. Often, the interest lies in comparing the transcriptional output between different conditions, and most RNA-seq studies can be classified as either: 1) differential gene expression (DGE) studies, where the overall transcriptional output of each gene is compared between conditions; 2) differential transcript/exon usage (DTU/DEU) studies, where the composition of a gene's isoform abundance spectrum is compared between conditions, or 3) differential transcript expression (DTE) studies, where the interest lies in whether individual transcripts show differential expression between conditions. DTE analysis results can be represented on the individual transcript level, or aggregated to the gene level, e.g., by evaluating

whether at least one of the isoforms shows evidence of differential abundance.

In this report, we make and give evidence for three claims: 1) gene-level estimation is considerably more stable than transcript-level; 2) regardless of the level at which abundance estimation is done, *inferences* at the gene level are appealing in terms of robustness, statistical performance and interpretation; 3) the magnitude of the difference between results obtained by simple counting and transcript-level abundance estimation is generally small in real data sets. However, despite strong overall correlations among results obtained from various quantification pipelines, taking advantage of transcript-level abundance estimates when defining or analyzing gene-level abundances leads to improved differential gene expression results compared to simple counting.

To facilitate a broad range of analysis choices, depending on the biological question of interest, we provide an R package, *tximport*, to import transcript lengths and abundance estimates from several popular quantification packages and export (estimated) count matrices and, optionally, average transcript length correction terms (i.e., offsets) that can be used as inputs to common statistical engines, such as $DESeq2^{11}$, $edgeR^{12}$ and $limma^{13}$.

Data

Throughout this manuscript, we utilize two simulated data sets and four experimental data sets (Bottomly¹⁴ [Data set 3], GSE64570¹⁵ [Data set 4], GSE69244¹⁶ [Data set 5], GSE72165¹⁷ [Data set 6], see Supplementary File 1 for further details) for illustration. Details on the data generation and full records of the analyses are provided in the data sets and Supplementary File 1. The first simulated data set (sim1; Data set 1) is the synthetic human data set from Soneson et al. 18, comprising 20,410 genes and 145,342 transcripts and is available from ArrayExpress (accession E-MTAB-3766). This data set has three biological replicates from each of two simulated conditions, and differential isoform usage was introduced for 1,000 genes by swapping the relative expression levels of the two most dominant isoforms. For each gene in this data set, the total transcriptional output is the same in the two conditions (i.e., no overall DGE); it is worth noting that this is an extreme situation, but provides a useful test set for contrasting DGE, DTU and DTE. The second simulated data set (sim2; Data set 2) is a synthetic data set comprising the 3,858 genes and 15,677 transcripts from the human chromosome 1. It is available from ArrayExpress with accession E-MTAB-4119. Also here, we simulated two conditions with three biological replicates each. For this data set, we simulated both overall DGE, where all transcripts of the affected gene showed the same fold change between the conditions (420 genes), differential transcript usage (DTU), where the total transcriptional output was kept constant but the relative contribution from the transcripts changed (420 genes) and differential transcript expression (DTE), where the expression of 10% of the transcripts of each affected gene was modified (422 genes, 528 transcripts). The three sets of modified genes were disjoint. Again, this synthetic data set represents an extreme situation compared to most real data sets, but provides a useful test case to identify underlying causes of differences between results from various analysis pipelines.

Data set 1.

http://dx.doi.org/10.5256/f1000research.7563.d109328

Data set 1 (html) contains all the R code that was used to perform the analyses and generate the figures for the **sim1** data set²⁸.

Data set 2.

http://dx.doi.org/10.5256/f1000research.7563.d109329

Data set 2 (html) contains all the R code that was used to perform the analyses and generate the figures for the **sim2** data set²⁹.

Data set 3.

http://dx.doi.org/10.5256/f1000research.7563.d109330

Data set 3 (html) contains all the R code that was used to perform the analyses and generate the figures for the Bottomly data set³⁰.

Data set 4.

http://dx.doi.org/10.5256/f1000research.7563.d109331

Data set 4 (html) contains all the R code that was used to perform the analyses and generate the figures for the **GSE64570** data set³¹.

Data set 5.

http://dx.doi.org/10.5256/f1000research.7563.d109332

Data set 5 (html) contains all the R code that was used to perform the analyses and generate the figures for the **GSE69244** data set²².

Data set 6.

http://dx.doi.org/10.5256/f1000research.7563.d109333

Data set 6 (html) contain all the R code that was used to perform the analyses and generate the figures for the ${\bf GSE72165}$ data set 3.

Gene abundance estimates are more accurate than transcript abundance estimates

To evaluate the accuracy of abundance estimation with transcript and gene resolution, we used *Salmon*⁷ (v0.5.1) to estimate TPM values for each transcript in each of the data sets. Gene-level TPM estimates, representing the overall transcriptional output of each gene, were obtained by summing the corresponding transcript-level TPM estimates. For the two simulated data sets, the true underlying TPM of each feature is known and we can thus evaluate the accuracy of the estimates. Unsurprisingly, gene-level estimates were more accurate than transcript-level estimates (Figure 1A, Supplementary Figures 1,2). We also derived TPM estimates from gene-level counts

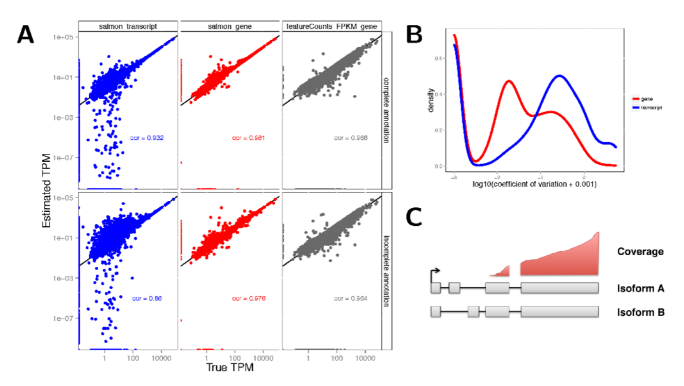


Figure 1 (sim2). A: Accuracy of gene- and transcript-level TPM estimates from *Salmon* and scaled FPKM estimates derived from simple counts from *featureCounts*, in one of the simulated samples (sampleA1). Spearman correlations are indicated in the respective panels. Top row: using the complete annotation. Bottom row: using an incomplete annotation, with 20% of the transcripts randomly removed. Gene-level estimates are more accurate than transcript-level estimates. Gene-level estimates from *Salmon* are more accurate than those from *featureCounts*. B: Distribution of the coefficients of variation of gene- and transcript-level TPM estimates from *Salmon*, calculated across 30 bootstrap samples of one of the simulated samples (sampleA1). Gene-level TPM estimates are less variable than transcript-level estimates. C: An example of unidentifiable transcript-level estimates, as uneven coverage does not cover the critical regions that would determine the amount that each transcript is expressed, while gene-level estimation is still possible.

obtained from featureCounts by dividing each of these with a reasonable measure of the length of the gene (the length of the union of its exons) and the total number of mapped reads, and scaling the estimates to sum to 1 million. The simple count estimates showed a lower correlation with the true TPMs than the Salmon estimates, in line with previous observations¹⁹. However, simple counts tended to show a high degree of robustness against incompleteness of the annotation catalog, as evidenced from estimation errors after first removing (at random) 20% of the transcripts (Figure 1A); in contrast, Salmon transcript estimate accuracies deteriorated. From the bootstrap estimates generated by Salmon, we also estimated the coefficient of variation of the abundance estimates. The gene-level estimates showed considerably lower variability in both simulated and experimental data (Figure 1B, Supplementary Figures 3,4). Taken together, these observations suggest that the gene-level estimates are more accurate than transcript-level estimates and therefore potentially allow a more accurate and stable statistical analysis. A further argument in favor of gene-level analysis is the unidentifiability of transcript expression that can result from uneven coverage caused by underlying technical biases (Figure 1C). Intermediate approaches, grouping together "indistinguishable" features are also conceiveable²⁰, but not yet standard practice.

DTE is more powerful and easier to interpret on gene level than for individual transcripts

DTE is concerned with inference of changes in abundance at transcript resolution, and thus invokes a statistical test for each transcript. We argue that this can lead to several complications: the first is conceptual, since the rows (transcripts) in the result table will in many cases not be interpreted independently, but will rather be grouping transcripts from the same gene, and the second one is more

technical, since the number of transcripts is considerably larger than the number of genes, which could lead to lower power due to the portioning of the total set of reads across a larger number of features and a potentially higher multiple testing penalty. We tested for DTE on the simulated data by applying $edgeR^{12}$ to the transcript counts obtained from Salmon (the application of count models to estimated counts is discussed in the next Section), and represented the results as transcript-level p-values or aggregated these to the gene level by using the perGeneQValue function from the DEXSeq²¹ R package. The transcript-level DTE test assesses the null hypothesis that the individual transcript does not change its expression, whereas the gene-level DTE test assesses the null hypothesis that all transcripts exhibit no change in expression. Framing the DTE question at the gene level results in higher power, without sacrificing false discovery rate control (Figure 2A). We note that this type of genelevel aggregation may favor genes in which one transcript shows strong changes, and that other approaches to increase power against specific alternatives are conceivable, e.g., capitalizing on the rich collection of methods for gene set analysis.

While DTE analysis is more suitable than DGE analysis for detecting genes with changes in absolute or relative isoform expression but no or only minor change in overall output (Supplementary Figure 5), we argue that even gene-level DTE results may suffer from lack of interpretability. DTE can arise in several different ways, from an overall differential expression of the gene or from differential relative usage of its transcripts, or a combination of the two (Figure 2B). We argue that the biological question of interest is in many cases more readily interpretable as a combination of DGE and DTU, rather than DTE. It has been our experience that results reported at the transcript level are still often cast to the gene level

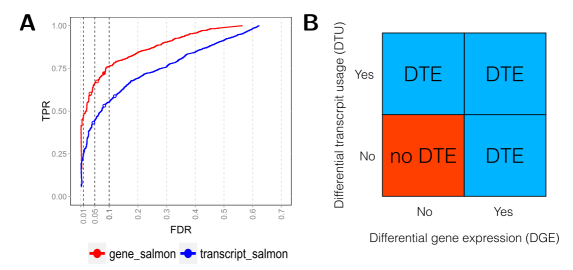


Figure 2 (sim2). A: DTE detection performance on transcript- and gene-level, using edgeR applied to transcript-level estimated counts from Salmon. The statistical analysis was performed on transcript level and aggregated for each gene using the perGeneQValue function from the DEXSeq R package; aggregated results show higher detection power. The curves trace out the observed FDR and TPR for each significance cutoff value. The three circles mark the performance at adjusted p-value cutoffs of 0.01, 0.05 and 0.1. B: Schematic illustration of different ways in which differential transcript expression (DTE) can arise, in terms of absence or presence of differential gene expression (DGE) and differential transcript usage (DTU).

(i.e., given a differentially expressed transcript, researchers want to know whether other isoforms of the gene are changing), suggesting that asking two specific gene-level questions (Is the overall abundance changing? Are the isoform abundances changing proportionally?) trumps the interpretability of one broad question at the transcript-level inference (Are there changes in any of the transcript expression levels?). Despite this, there are of course also situations when a transcript-centric approach is superior, for example in targeted experiments where specific isoforms are expected to change due to an administered treatment.

Incorporating transcript-level estimates leads to more accurate DGE results

DGE (i.e., testing for changes in the overall transcriptional output of a gene) is typically performed by applying a count-based inference method from statistical packages such as $edgeR^{12}$ or $DESeq2^{11}$ to gene counts obtained by read counting software such as featureCounts¹, HTSeq-count² or functions from the GenomicAlignments²² R package. A lot has been written about how simple counting approaches are prone to give erroneous results for genes with changes in relative isoform usage, due to the direct dependence of the observed read count on the transcript length²³. However, the extent of the problem in real data has not been thoroughly investigated. Here, we show that taking advantage of transcript-resolution estimates (e.g., obtained by Salmon) can lead to improved DGE results. We propose two alternative ways of integrating transcript abundance estimates into the DGE pipeline: to define an "artificial" count matrix, or to calculate offsets that can be used in the statistical modeling of the observed gene counts from, e.g., featureCounts. Both approaches are implemented in the accompanying tximport R package (available from https://github.com/mikelove/tximport).

We defined three different count matrices for each data set: 1) using *featureCounts* from the *Rsubread*¹ R package (denoted **featureCounts** below), 2) summing the estimated transcript counts from *Salmon* within genes (**simplesum**), 3) summing the estimated transcript TPMs from *Salmon* within genes, and multiplying with the total library size in millions (**scaledTPM**). We note that the scaledTPM values are artificial values, transforming underlying abundance measures to the "count scale" to incorporate the information provided by the sequencing depth. We further used the *Salmon* transcript lengths and estimated TPMs to define average transcript lengths for each gene and each sample (normalization factors) as described in the Supplementary material, to be used as offsets for *edgeR* and *DESeq2* when analyzing the feature-Counts and simplesum count matrices (**featureCounts_avetxl** and **simplesum_avetxl**).

Overall, the counts obtained by all methods were highly correlated (Supplementary Figures 6–8), which is not surprising since any differences are likely to affect a relatively small subset of the genes. In general, the simplesum and featureCounts matrices led to similar conclusions in all considered data sets. However, there are differences between the two approaches in terms of how multimapping reads and reads partly overlapping intronic regions are handled²⁴. The concordance between simplesum and featureCounts results also suggests that statistical methods based on the Negative Binomial assumption are applicable also to summarized, gene-level estimated counts, which is further supported by the similarity

between the p-value histograms as well as the mean-variance relationships observed with the three types of count matrices (Supplementary Figures 9–14).

Accounting for the potentially varying average transcript length across samples when performing DGE, either in the definition of the count matrix (scaledTPM) or by defining offsets, led to considerably improved false discovery rate (FDR) control compared to using the observed featureCounts or aggregated Salmon counts (simplesum) directly (Figure 3A, Table 1). It is important to note that this improvement is entirely attributable to an improved handling of genes with changes in isoform composition between the conditions (Figure 3B, Supplementary Figure 15), that we purposely introduced strong signals in the simulated data set in order to pinpoint these underlying causes, and that the overall effect in a real data set will depend on the extent to which considerable DTU is present. Experiments on various real data sets (Supplementary Figure 16) show only small differences in the collections of significant genes found with the simplesum and simplesum_avetxl approaches, suggesting that the extent of the problem in many real data sets is limited, and that most findings obtained with simple counting are not induced by counting artifacts. Further support for this conclusion is shown in Figure 4 (see also Supplementary Figures 17–19 and Supplementary Table 1), where log-fold change estimates from edgeR, based on the simplesum and scaledTPM matrices, are contrasted. For the genes with induced DTU in the sim2 data set, log-fold changes based on the simplesum matrix are overestimated, as expected. However, this effect is almost absent in all the real data sets, again highlighting the extreme nature of our simulated data and suggesting that the effect of using different count matrices is considerably smaller for many real data sets. Table 1 suggests that the lack of error control for simplesum and featureCounts matrices is more pronounced when there is a large difference in length between the differentially used isoforms. In the group with smallest length difference, where the longer differentially used isoform is less than 34% longer than the shorter one, all approaches controlled the type I error satisfactorily. It is worth noting that among all human transcript pairs in which both transcripts belong to the same gene, the median length ratio is 1.85, and for one third of such pairs the longer isoform is less than 38% longer than the shorter one (see Data set 1).

Discussion

In this article, we have contrasted transcript- and gene-resolution abundance estimation and statistical inference, and illustrated that gene-level results are more accurate, powerful and interpretable than transcript-level results. Not surprisingly, however, accurate transcript-level estimation and inference plays an important role in deriving appropriate gene-level results, and it is therefore imperative to continue improving abundance estimation and inference methods applicable to individual transcripts, since misestimation can propagate to the gene level. We have shown that when testing for changes in overall gene expression (DGE), traditional gene counting approaches may lead to an inflated false discovery rate compared to methods aggregating transcript-level TPM values or incorporating correction factors derived from these, for genes where the relative isoform usage differs between the compared conditions. These correction factors can be calculated from the output of transcript abundance programs, using e.g., the provided R package (tximport). It is important to note that the average transcript length

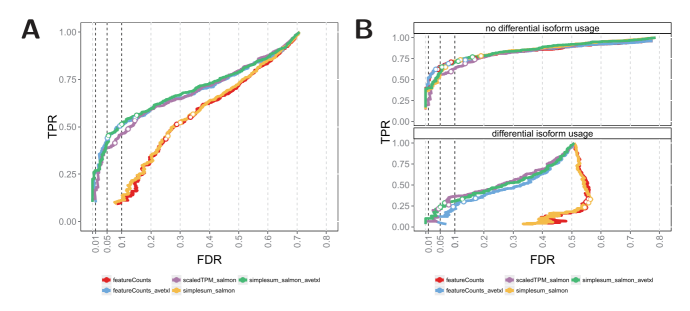


Figure 3 (sim2). A: DGE detection performance of *edgeR* applied to three different count matrices (simplesum, scaledTPM, featureCounts), with or without including an offset representing the average transcript length (for simplesum and featureCounts). Including the offset or using the scaledTPM count matrix leads to improved FDR control compared to using simplesum or featureCounts matrices without offset. The curves trace out the observed FDR and TPR for each significance cutoff value. The three circles mark the performance at adjusted p-value cutoffs of 0.01, 0.05 and 0.1. **B**: stratification of the results in **A** by the presence of differential isoform usage. The improvement in FDR control seen in **A** results from an improved treatment of genes with differential isoform usage, while all methods perform similarly for genes without differential isoform usage.

Table 1 (sim1). Observed false positive rates from a differential gene expression analysis using edgeR applied to various count matrices (with a nominal p-value cutoff at 0.05), limited to genes with true underlying differential isoform usage (recall that no genes are truly differentially expressed in this data set). The results are stratified by "effect size" (the difference in relative abundance between the two differentially used isoforms) and the length ratio between the longer and the shorter of the differentially used isoforms. FPRs below the nominal p-value threshold (0.05) are marked in bold. For more details, see Data set 1.

	simplesum	featureCounts	simplesum_avetxI	featureCounts_avetxl	scaledTPM
[0,0.33], [1,1.34]	0.019	0.019	0.023	0.023	0.023
[0.33,0.67], [1,1.34]	0.059	0.059	0.059	0.059	0.059
[0.67,1], [1,1.34]	0.000	0.053	0.053	0.053	0.053
[0,0.33], [1.34,2.57]	0.075	0.070	0.070	0.065	0.065
[0.33,0.67], [1.34,2.57]	0.240	0.220	0.050	0.033	0.066
[0.67,1], [1.34,2.57]	0.420	0.540	0.038	0.077	0.038
[0,0.33], [2.57,35.4]	0.150	0.140	0.037	0.043	0.037
[0.33,0.67], [2.57,35.4]	0.650	0.650	0.060	0.060	0.034
[0.67,1], [2.57,35.4]	0.970	0.970	0.034	0.034	0.034

offsets must account for the differences in transcript usage between the samples and thus using (sample-independent) exon-union gene lengths will not improve performance.

All evaluated counting approaches gave comparable results for genes where DTU was not present. Thus, the extent of the FDR inflation in experimental data depends on the extent of DTU between the compared conditions; notably, our simulation introduced rather extreme levels of DTU, hence the inflated FDR, and

the difference between the approaches was considerably smaller in real data sets. Recent studies have also shown that many genes express mainly one, dominant isoform²⁵ and for such genes, we expect that simple gene counting will work well.

Our results highlight the importance of correctly specifying the question of interest before selecting a statistical approach. Summarization of abundance estimates at the gene level before performing the statistical testing should be the method of choice if the

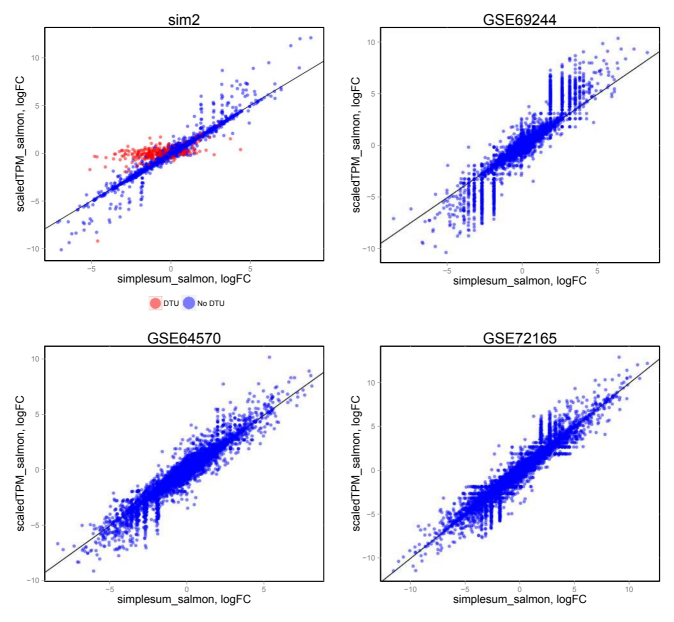


Figure 4. Comparison of log-fold change estimates from edgeR, based on simplesum and scaledTPM count matrices, in four different data sets. For the simulated data set (sim2), where signals have been exaggerated to pinpoint underlying causes of various observations, genes with induced DTU (whose true overall log-fold change is 0) show a clear overestimation of log-fold changes when using simplesum counts. However, none of the real data sets contain a similar population of genes, suggesting that for many real data sets, simple gene counting leads to overall similar conclusions as accounting for underlying changes in transcript usage.

interest is in finding changes in the overall transcriptional output of a gene. However, it is suboptimal if the goal is to identify genes for which *at least one* of the transcripts show differences in transcriptional output, since it may miss genes where two transcripts change in opposite directions, or where a lowly expressed transcript changes. For gene-level detection of DTE (that is, whether any transcript showed a change in expression between the conditions), statistical testing applied to aggregated gene counts led to reduced power and slightly inflated FDR compared to performing the

statistical test on the transcript level and aggregating results within genes (Supplementary Figure 5). Statistical inference on aggregated transcript TPMs (scaledTPM) showed low power for detecting changes that did not affect the overall transcriptional output of the gene, as expected. An alternative to DTE analysis, for potential improved interpretability, is to perform a combination of DGE and DTU analyses, both resulting in gene-level inferences. Table 2 summarizes our results and give suggested workflows for the different types of analyses we have considered.

Table 2. Summary of suitable analysis approaches for the three types of comparative analyses discussed in the manuscript (DGE, DTE and DTU).

Task	Input data	Software (examples)	Post-processing	
DGE	Aggregated transcript counts + average transcript length offsets, or	Salmon, kallisto, BitSeq, RSEM		
	simple counts + average transcript length offsets	tximport		
		DESeq2, edgeR, voom/ limma		
DTE	Transcript counts	Salmon, kallisto, BitSeq, RSEM	Optional gene-level aggregation	
		tximport		
		DESeq2, edgeR, sleuth, voom/limma		
DTU/DEU	Transcript counts or bin counts, depending on interpretation potential ¹⁸	Salmon, kallisto, BitSeq, RSEM	Optional gene-level aggregation	
		DEXSeq		

Of course, there may be situations where a direct transcript-level analysis is appropriate. For example, in a cancer setting where a specific deleterious splice variant is of interest (e.g., AR-V7 in prostate cancer²⁶), inferences directly at the transcript level may be preferred. However, while this may be preferred for individual known transcripts, transcriptome-wide differential expression analyses may not be warranted, given the associated multiple testing cost.

Finally, we note that estimation at the gene level can reduce the problem of technical biases on expression levels and unidentifiable estimation. Current methods for transcript-level quantification (e.g., *Cufflinks, RSEM, Salmon, kallisto*) do not correct for amplification bias on fragments, which can lead to many estimation errors, such as expression being attributed to the wrong isoform²⁷. Non-uniform coverage from amplification bias or from position bias (3' coverage bias from poly-(A) selection) can result in unidentifiable transcript-level estimation. Such errors and estimation problems are minimized when summarizing expression to the gene level.

Data availability

F1000Research: Data set 1. 10.5256/f1000research.7563.d109328

F1000Research: Data set 2. 10.5256/f1000research.7563.d109329

F1000Research: Data set 3. 10.5256/f1000research.7563.d109330

F1000Research: Data set 4. 10.5256/f1000research.7563.d109331

F1000Research: Data set 5. 10.5256/f1000research.7563.d109332

F1000Research: Data set 6. 10.5256/f1000research.7563.d109333

Software availability

Software access

https://github.com/mikelove/tximport

Source code as at the time of publication

https://github.com/F1000Research/tximport

Archived source code as at the time of publication http://dx.doi.org/10.5281/Zenodo.35123

Software license

tximport is released under a GNU Public License (GPL).

Author contributions

CS, MIL and MDR conceived the study and developed methodology. CS and MDR designed and carried out the computational experiments and drafted the manuscript. MIL implemented the *tximport* R package and wrote parts of the manuscript. All authors read and approved the final manuscript and have agreed to the content.

Competing interests

No competing interests were disclosed.

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Supplementary material

Supplementary File 1

Supplementary File 1 (pdf) contains more detailed information about the data sets, supplementary methods and supplementary figures referred to in the text.

Click here to access the data.

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Rob Patro 🗓

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In this manuscript, the authors address a few questions of considerable (and perennial) interest in the analysis of RNA-seq data. Specifically, they provide evidence that, using available methods (e.g. DESeq2 / edgeR), assessing differential expression at the gene-level (DGE) is more robust than at the transcript level (DTE). Further, they convincingly argue that estimating abundance at the level of transcripts, and then aggregating these abundances to the gene level leads to improved estimation of differential gene expression. They demonstrate that one of the major factors in this improved estimation is the availability of a sample-specific feature length for each gene (derived from the abundance-weighted length of the expressed transcripts of this gene), which is not possible to obtain with any fixed gene model used by count-based methods. Finally, the authors argue that much of the analysis of interest at the transcript level does not actually require differential transcript expression testing, but rather can be inferred from a combination of DGE and differential transcript usage (DTU); this is an interesting proposition that merits further discussion and analysis. Overall, this is a well-written paper, with extensive and compelling supplementary and supporting data, that addresses a ubiquitous analysis task involving RNA-seg. It should be of broad interest to the community and makes a valuable contribution. The accompanying software, tximport, is user-friendly and makes it easy to apply the type of analysis recommended herein; it too should be widely useful.

Major comments:

It would be very useful to provide the equations used for calculating each of the abundance measured considered directly. Section 4 of the supplementary information is useful to this end, but the reader still has to search a bit to see exactly how each metric is computed (though the fantastic R-Markdown included with the figures means that these computations can be found explicitly).

Similarly, it would be useful to the reader to provide a description, in prose, of how specific experiments were performed (again, the reproducible nature of most of these experiments makes tracking down this information possible, but sometimes time-consuming). For example, how,

precisely, was removal of transcripts handled at the level of the genome annotation? If a transcript consists only of constitutive exons, were all of those exons retained in the genome annotation used for STAR + featureCounts, while the transcript was removed in the Salmon index?

The result that transcript-level abundance estimation is more sensitive to the removal of transcripts than gene-level abundance estimation — this seems intuitive. However, I agree with Dr. Floor's suggestion that:

"The assertion that "simple counts tended to show a high degree of robustness against incompleteness of the annotation catalog, as evidenced from estimation errors after first removing (at random) 20% of the transcripts" seems misleading since Salmon-derived gene-level abundances actually show a higher Spearman correlation than count-derived gene-level abundances when subjected to removing a random 20% of transcripts."

I would suggest rewording this sentence, as the main result seems to be that gene-level analysis is more robust to an incomplete annotation than transcript-level analysis. Transcript-level abundance estimation followed by gene-level analysis seems to perform just as well (actually, better) than gene-level counting in this scenario.

The experiments in the section "Incorporating transcript-level estimates leads to more accurate DGE results" suggests the (reasonable) interpretation that the main benefit of incorporating transcript-level abundance estimates when assessing DGE is a more accurate measure of the "feature" length of the gene. The authors state "It is important to note that this improvement is entirely attributable to an improved handling of genes with changes in isoform composition between the conditions." This is supported by the fact that using the abundance-weighted average transcript length (i.e. offsets) with counting based approaches improves the results substantially. However, one other place where transcript-level abundance estimates are useful in the context of DGE is when assessing the expression of paralogous genes. While most multi-mapping reads that derive from different isoforms of the same gene will be uniquely mappable at the level of the genome, and hence will be included in the counts for that gene, reads that map ambiguously among paralogs may not be. In such cases, count-based methods do not have a principled way of apportioning a read between the paralogs involved, and discarding multi-mapping reads may negatively affect estimation of the abundance of the paralogs, even at the gene level. While this case is likely much less common than mis-estimation of DGE as a result of DTU, it is certainly of biological interest. I would suggest adding an analysis, restricted to sets of paralogous genes, comparing how the different approaches perform in this case. This may help to highlight the importance of not only deriving appropriately weighted and sample-dependent lengths for genes, but also on resolving multi-mapping ambiguity that occurs between genomically distinct loci.

The argument that most transcript-level analyses of interest may be addressed by looking at DGE in conjunction with DTU is an interesting one. It is certainly that case that not all tasks for which DTE is used actually require assessing differential expression at the transcript level. One issue with the DGE + DTU-based analysis which warrants further discussion in the manuscript is that I believe that this approach, too, would require correcting for multiple hypothesis testing. Specifically, one is testing both the DGE and the DTU hypotheses for each gene (or for a relevant subset of interest). The correction here is likely to be less harsh than in the case of assessing DTE, but is still worth discussing.

Minor comments:

As per Dr. Floor's statement, Salmon (and Sailfish) also incorporate sequence-specific bias correction. Further, RSEM and Salmon (and a few other transcript-level abundance estimation tools) also incorporate the modeling of non-uniform fragment start position distributions. Of course, modeling a non-uniform start position distribution cannot overcome a complete lack of sampling in critical regions that might make determining transcript-level fragment assignment impossible, but it may help in properly apportioning an ambiguously-mapped fragment between transcripts depending on its relative position in each.

One potential added source of variability here is that all Salmon estimates presented in the manuscript make use of Salmon's quasi-mapping of reads, while the STAR + featureCount pipeline makes use of "traditional" alignments. This is the primary intended usage mode of Salmon, and absolutely does represent a "typical" pipeline for methods that avoid alignment (Salmon, Sailfish, kallisto). However, it would probably be best to mention this as a potential (though likely negligible) additional source of variability.

In the discussion, the authors argue that "... it is therefore imperative to continue improving abundance estimation and inference methods applicable to individual transcripts, since misestimation can propagate to the gene level." This is, of course, an important and valid suggestion. Another direction, on which it would be useful to get the authors' thoughts and suggestions, is the development of differential expression tools (at either the transcript or gene level) that can make use of the variance estimates that some tools (like Salmon) can provide. To what extent might incorporating this information help control false positive rates and improve DTE or even DGE estimates?

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 19 Feb 2016

Mark Robinson, University of Zurich, Switzerland

Thank you for taking the time to read and review our paper. As per your suggestions, we have expanded the supplementary pdf document with equations defining each of the abundance measures that we used as well as a detailed description of the generation of the incomplete annotation files. We have also reworded the paragraph discussing the influence of an incomplete annotation catalog, and added further discussion points regarding multiple testing correction for DGE+DTU as well as regarding new methods incorporating variance estimates into the differential analysis. However, a deeper discussion of the latter point falls outside the scope of this article.

We found your suggestion to restrict the abundance accuracy comparison to paralogous genes interesting, and the supplement has been expanded with several examples comparing the accuracy of gene-level abundance estimates from Salmon and from STAR+featureCounts, restricted to sets of paralogous genes. While we see a clear advantage

of Salmon for many of the paralogous gene groups, the overall difference is only slightly larger for sets of paralogous genes compared to random sets of genes (as can be seen in the revised version of Dataset 1). In addition, we have added the following text to the relevant section in the results:

"...suggesting that the extent of the problem in many real data sets is limited... though some loss of sensitivity for certain genes may be encountered from discarding multimapping fragments, which may be recovered through the use of transcript abundance estimators such as Salmon."

While we acknowledge that many of the popular transcript abundance estimation methods incorporate some type of bias estimation/correction (albeit not at the fragment level), and that it is definitely an important (and difficult) research area, a thorough discussion on the relative merits of different bias correction approaches is outside the scope of this paper. We have added citations to relevant literature on coverage bias, have simplified and clarified the last paragraph of the discussion, and have reworded the sentence about unidentifiable estimation due to coverage biases:

"While some extent of coverage variability might be alleviated by corrections for sequenceor position-specific biases, there remain cases where transcript expression cannot be inferred from data (Figure 1C)"

Competing Interests: No competing interests were disclosed.

Reviewer Report 04 January 2016

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Stephen N. Floor 🗓

Department of Molecular and Cell Biology, University of California, Berkeley, Berkeley, CA, USA

Soneson, Love and Robinson tackle a crucial question for analysis of RNA deep sequencing data in this manuscript: what is the role of transcript diversity in the accuracy and statistical power associated with measurements of gene expression? The authors make and convincingly show three claims: gene-level estimation and inferences are more robust than those at the transcript-level, and incorporating transcript-level quantification into gene-level abundance leads to improved differential expression testing. The claims are convincingly proven, the manuscript is well written, and the subject matter is of considerable interest. Furthermore, the described R package *tximport* should be of broad interest to the RNA deep sequencing community.

Overall comments:

It may be useful to indicate explicitly in the text that the methods are contained within the (excellently written and formatted) supplementary material, as this was not apparent. It might be clearest to create a specific methods section that just references supplementary file 1.

The clarity of scatter plots with more than ~hundreds of points (e.g. Figure 1A) could be improved by using partially transparent points to visualize density.

Introduction:

Paragraph 1: Cufflinks, RSEM and Bitseq are grouped with kallisto and Salmon and it is then stated that some of these methods bypass read alignment. It would be clearer if this were reworded to avoid the ambiguity as to which methods avoid read alignment.

Paragraph 4: The third claim could be presented more clearly. While it is interesting that simple counting performs similarly to transcript-level quantification procedures, it seems more interesting to this reviewer that incorporating transcript-level information improves the accuracy of differential expression testing at the gene level. Perhaps these two concepts can be combined into one more concise point?

Results:

The assertion that "simple counts tended to show a high degree of robustness against incompleteness of the annotation catalog, as evidenced from estimation errors after first removing (at random) 20% of the transcripts" seems misleading since Salmon-derived gene-level abundances actually show a higher Spearman correlation than count-derived gene-level abundances when subjected to removing a random 20% of transcripts. Figure 1a bottom left shows that transcript-level abundances are strongly affected by removal of 20% of transcripts, but that gene-level abundances are not strongly changed whether estimated using counts or Salmon. This statement should be reworded.

Two concerns are raised about DTE. It is certainly true that reads are spread across more features when performing DTE as opposed to DGE. However, it is not apparent why analysis of DTE involves grouping of transcripts together for interpretation. DTE implies analysis at the transcript level and therefore no grouping, while DGE could involve some level of grouping of transcripts or quantification at the gene level from the start. The clarity of this could be improved.

It is a very interesting idea to separately frame questions regarding DGE and DTU, which should be adopted widely, as the two are separable questions.

The authors state one possible workflow towards DGE analysis in the section "Incorporating transcript-level estimates leads to more accurate DGE results." Alternative pipelines (e.g. cuffdiff) could be presented in brief.

The observation that simplesum and featureCounts results are highly correlated and therefore that statistical methods based on the Negative Binomial distribution can be used on estimated counts seems of greater importance than is emphasized in the text. This should be elaborated upon in the discussion, since this means that estimated counts from kallisto, express, salmon, etc

can be used directly by statistical packages assuming a NB distribution (edgeR, DESeq2, etc). This point is frequently debated in discussions of how to rigorously analyze sequencing data. The conclusion here that NB applies to estimated counts is thus quite important.

Please explain the meaning of the name for each curve in the legend for Figure 3 (i.e. specify that "avetxl" means using the offset corresponding to average transcript length.

Discussion:

The assertion that "gene-level results are more accurate, powerful and interpretable than transcript-level results" seems an oversimplification given the result that incorporating transcript-level quantification leads to improved DGE detection performance (e.g. Fig 3).

Please cite at minimum Roberts *et al.*, (2011) regarding sequence bias correction as this has been implemented in cufflinks, express and kallisto. Other relevant papers should also be included here, as attempts have been made to address both positional and sequence-specific bias in RNA sequencing data.

Supplement:

The usability of the supplemental info could be improved by substituting rasterized for vectorized plots for those with ~hundreds of points.

Please explain the meaning of the name for each curve in the legend for Supplemental Figure 5.

References

1. Roberts A, Trapnell C, Donaghey J, Rinn JL, et al.: Improving RNA-Seq expression estimates by correcting for fragment bias. *Genome Biol.* 2011; **12** (3): R22 PubMed Abstract | Publisher Full Text

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 19 Feb 2016

Mark Robinson, University of Zurich, Switzerland

Thank you for taking the time to read and review our paper. In the revised version, we have improved the clarity and usability of the figures and the supplement by using partially transparent points and extending the figure captions. We have also modified the main text to refer more clearly to the method information provided in the supplementary material, and improved the clarity of the text in several places, according to your suggestions.

For additional comments regarding references to bias correction studies, please see responses to Rob Patro.

Competing Interests: No competing interests were disclosed.

Comments on this article

Version 1

Author Response 02 Feb 2016

Charlotte Soneson, University of Zurich, Zurich, Switzerland

Hi Nick,

thanks for your comments.

Regarding the increased accuracy and robustness of gene-level estimates compared to transcript-level ones (even with the full annotation, bootstrap variances of gene estimates are lower than those of transcript estimates), we agree that it is not surprising. However, we still found it worth pointing out since it implies that in some situations, gene-level statistical analyses may be preferred due to the increased precision of the input data.

For your second point, note that Figure 2 does not in fact compare DGE to DTE, but rather DTE summarized on gene- and transcript-level. You are of course right that just reducing the number of tests does not automatically result in higher power. Instead, the difference between the gene- and transcript-level results in Figure 2 is largely due to the different null hypotheses (as outlined in the text) and, consequently, the type of signal we require to call a feature (gene or transcript) significant. In fact, the gene-level summarized analysis is answering a somewhat "easier" question, from the sensitivity point of view. Consider for example a gene with 5 differentially expressed transcripts. On the gene level, we can reach a power of 100% for this gene if **any** of the transcripts is considered significantly DE. On the transcript level, we would need all of the transcripts to be significant to reach the same power. Looking at this particular data set, for many of the genes that are found as true positives with the gene level test, not all truly DE transcripts are actually detected. This explains the lower power of the transcript-level analysis. We will clarify this in the revised version. Regarding the choice of DTE vs DGE+DTU, it is clear that one solution will not always be the optimal choice, and it will likely depend on the particular problem as well as on the person interpreting the results. However, we have found that in many of our own collaborations, the biological question can be more clearly stated in terms of DGE+DTU (not necessarily performed sequentially, rather in parallel). Part of the reason for this discussion was to encourage researchers to think about what question they really want to answer before starting their analyses.

Finally, note that the only place where we are actually comparing "summarizing DTE at the gene level" and DGE is in Supplementary Figure 5 (on simulated data). In Figures 3-4, the question of interest is always comparing the total transcriptional output of a gene between conditions (i.e., DGE), and all methods are based on aggregating gene abundance estimates before the statistical

test is applied. As you note, we don't see a big global effect of including offsets accounting for average transcript lengths for real data. However, there may of course still be important effects for individual genes, where isoforms of different lengths are expressed in different conditions.

Thanks again for your comments, and we are glad you found the paper useful!

Competing Interests: No competing interests were disclosed.

Reader Comment () 29 Jan 2016

Nick Schurch, The Barton Group, Division of Computational Biology College of Life Sciences, University of Dundee, UK

I read this work with great interest for a literature review and found it engaging and informative. I thought I'd post my resulting thoughts here.

This paper focusses on the advantages and disadvantages of performing differential expression analysis at either the gene level or the transcript level, relying heavily on simulated data. Initially the authors use simulated data to show that gene level estimations of expression are more stable then transcript level estimations. They do this by showing that with the removal of ~20% of the transcripts from the annotation, the quantification the transcript level estimations becomes less accurate while the gene level estimation remains relatively unchanged. I found this conclusion to be unsurprising; the transcripts are the primary annotation used for transcript level quantification but are not the primary annotation for the gene level estimation. Except in rare cases where the removal of the transcripts result in the removal of the parent gene or removal of the longest transcripts results in a considerable change in the length of the parent gene, the gene annotation will remain largely unaltered by the removal of transcripts.

The authors then consider the performance of differential expression at the gene level (DGE) vs differential expression at the transcript level (DTE), concluding that inference at the gene level is more robust, easier to interpret and has better statistical performance. While the argument for improved robustness is compelling, I found the arguments for improved inference and statistical performance a little misleading. The improved statistical performance of DGE is due essentially to performing fewer tests. By this argument, anything that reduced the number of tests always improves the inference (e.g., using a smaller annotation) which is clearly not always the case. The authors also suggest that rather than identifying DTE, a better approach with information summarized at the gene level may be to first identify which genes are changing and then identify differential isoform usage. There is no clear demonstration of the advantages of this approach or what the appropriate multiple testing correction is for the combined analysis. In particular, I find the interpretability of DTE far more straightforward than the combined approach.

The paper also examined the differences between the differential gene expression conclusions resulting from summarizing DTE at the gene level and DGE called on simple gene counts on both simulated and real data and find that, again somewhat unsurprisingly in my opinion, that the difference in the results is smaller and less obvious for real datasets than for the simulated data

where they have added a strong signal. I found the evidence for using an adjusted transcript length is compelling despite the relatively small apparent impact it has for real datasets.

All in all I liked the paper and it was interesting to talk about it and discuss the issues it raises with my group.

Competing Interests: No competing interests were disclosed.

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