Thousands of exon skipping events differentiate among splicing patterns in sixteen human tissues [version 1; referees: 2 approved]

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Abstract
Alternative splicing is widely recognized for its roles in regulating genes and creating gene diversity. However, despite many efforts, the repertoire of gene splicing variation is still incompletely characterized, even in humans. Here we describe a new computational system, ASprofile, and its application to RNA-seq data from Illumina’s Human Body Map project (>2.5 billion reads). Using the system, we identified putative alternative splicing events in 16 different human tissues, which provide a dynamic picture of splicing variation across the tissues. We detected 26,989 potential exon skipping events representing differences in splicing patterns among the tissues. A large proportion of the events (>60%) were novel, involving new exons (~3000), new introns (~16000), or both. When tracing these events across the sixteen tissues, only a small number (4-7%) appeared to be differentially expressed (‘switched’) between two tissues, while 30-45% showed little variation, and the remaining 50-65% were not present in one or both tissues compared. Novel exon skipping events appeared to be slightly less variable than known events, but were more tissue-specific. Our study represents the first effort to build a comprehensive catalog of alternative splicing in normal human tissues from RNA-seq data, while providing insights into the role of alternative splicing in shaping tissue transcriptome differences. The catalog of events and the ASprofile software are freely available from the Zenodo repository (http://zenodo.org/record/7068; doi:10.5281/zenodo.7068) and from our website http://ccb.jhu.edu/software/ASprofile.

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Background

Alternative splicing is a widespread phenomenon in eukaryotic species, and differential regulation of alternative splice variants is gaining recognition as an important mechanism of gene regulation. More than 90% of human genes are estimated to be alternatively spliced\(^1\), producing multiple transcripts and (often) different protein sequences from a single locus. The number of variants of a gene ranges from two to potentially thousands\(^1\). The resulting proteins may exhibit different and sometimes antagonistic functional and structural properties\(^1\), and may inhabit the same cell with the resulting phenotype representing a balance between their expression levels\(^2\). Defects in splicing have been implicated in human diseases, including cancer\(^3-6\). Developing a comprehensive catalog of splice variant annotations across a wide range of tissues and conditions is important not only as part of our efforts to create a complete gene list for the human genome, but also to serve as a reference for differential expression studies aiming to identify molecular markers of disease.

Annotation of alternative splicing has traditionally been based on cDNA (expressed sequence tags (EST), mRNA) sequence data from public repositories such as dbEST, RefSeq\(^1\), and the Mammalian Gene Collection\(^1\). These data sources were compiled over many years, from independent contributions by thousands of investigators working on different genes and systems, and are therefore inconsistent in their coverage of the transcriptome in general and of each gene individually. Because these resources were generated using Sanger sequencing, they were relatively expensive to produce, but despite the cost have insufficient depth to capture the diversity of splicing variations in human cells. RNA-seq technology produces vastly more sequence data in a cost-effective way and in a much shorter amount of time, allowing a deep characterization of the transcriptome in a variety of cells and conditions\(^6-11\). However, even though the Illumina Human Body Map, we addressed the questions: “How much alternative splicing do we find?” and “How does alternative splicing vary among tissues?” We used this data set, spanning 16 tissues and containing over 2.5 billion sequences, to build a comprehensive catalog of alternative splicing (AS) within each tissue. We also compared AS profiles across tissue types to derive insights into the role of AS in shaping transcriptome differences.

Results

We analyzed the Illumina Human Body Map RNA-seq set (ArrayExpress accession: E-MTAB-513; http://www.ebi.ac.uk/arrayexpress), consisting of approximately 160 million reads from each of 16 tissues, each from a different individual. This resource is one of the most high-quality and complete to date, and therefore allows us to detect AS events with high accuracy. To determine splicing variations in each tissue, we first mapped reads to the reference genome and assembled them into transcripts or transcript fragments. Then we analyzed the transcripts to determine putative alternative splicing events, in particular exon skipping events, within and between samples, and compared them across the tissues. We focused on exon skipping because the alignment evidence for these events is usually clear and unambiguous, and less likely to be confounded by alignment or assembly artifacts. The data support a number of overall findings:

1. Based on a comparison against several annotation databases (Ensembl\(^13\), CCDS\(^16\), UCSC Genes\(^17\) and H-ASDB\(^19\)), we found that 11–45% of the assembled transcripts in each tissue were unannotated, as well as a majority (65%) of the 26,989 exon skipping events discovered from this data set.
2. These novel events appear to be more tissue-specific than previously annotated (known) events; i.e., they tend to occur in fewer tissue types.
3. When an exon is skipped, it usually occurs in a different tissue from those in which it is present; only 5–23% of events express both forms within the same tissue.
4. Comparing exon skipping profiles across tissues, we found that only 10–20% of the events identified show different splicing ratios between any two given tissues, whereas 50–65% of the cataloged events are not present in either or both tissues.

Overall, our analysis reveals a complex and dynamic picture of alternative splicing across tissue types, where differences among tissue transcriptomes arise from the interplay between constitutive transcription and alternative splicing. Most importantly, we compiled the first large repository of putative exon skipping and other classes of alternative splicing events in normal human tissues detected from RNA-seq data, which will be a valuable resource for studies of regulation and to identify markers of diseases. This catalog and our methods, implemented in the open source software program ASprofile, are freely available from the Zenodo repository (http://zenodo.org/record/7068; doi:10.5281/zenodo.7068) and from our web site http://ccb.jhu.edu/software/ASprofile.

A global view of alternative splicing in the 16 tissues

To determine alternative splicing events and globally characterize alternative splicing within a given tissue, we analyzed 50-bp paired-end sequences from 16 different tissues: adrenal, adipose, brain, breast, colon, heart, kidney, liver, lung, lymph, ovary, prostate, skeletal muscle, testis, thyroid and white blood cells. These data are publicly available as the Illumina Human Body Map project (EMBL accession ENA-ERP000546; ArrayExpress accession: E-MTAB-513; http://www.ebi.ac.uk/arrayexpress/browse.html?keywords=E-MTAB-513&expandefo=on). Libraries were made from polyA-selected mRNA with an insert size of 210 bp, independently for each tissue, using a random priming process and unstranded. One run of 2x50 bp paired-end sequencing was performed on the Illumina HiSeq2000 instrument, using one lane per tissue, to produce approximately 80 million pairs of reads (160 million sequences) per tissue. The entire data set comprises ~128 gigabases (GB) of sequence (~8 GB sequence per tissue), making this one of the most complete RNA-seq resources to date and one of very few spanning multiple types of tissues.

We mapped reads to the human genome with the program TopHat\(^15\) (Supplemental Table S1), and then assembled overlapping reads on the genome into transcript fragments using Cufflinks\(^18\). Cufflinks represents all reads at a locus as an assembly graph, in which any two reads are connected if they overlap and have compatible splice patterns, and
then traverses the graph to produce the minimum number of transcripts that can explain all of the input reads. Because single-exon transcripts, which form the bulk of the assemblies (Figure 1 and Supplemental Table S2), are frequently artifacts of sequencing and mapping, we used only the multi-exon transcripts to measure the gene and transcript content.

Although the assemblies produced by Cufflinks can be full-length transcripts, many transcripts can only be assembled into partial fragments (e.g., when the coverage of a transcript contains gaps). We therefore designate all transcript assemblies, complete or otherwise, as *transfrags*. For each tissue, Cufflinks produced between 23,000–46,000 multi-exon transfrags, clustered into 20,000–30,000 loci. The number of transfrags was greatest in brain and testes, and lowest in liver, colon, white blood cells and skeletal muscle, reflecting the combined effects of the number of expressed genes (Pearson’s $r^2=0.74$) and splicing variation within genes. These findings are consistent with some of the earlier estimates of the sizes of transcriptomes of different tissues\(^1\),\(^2\)–\(^25\). To estimate the number of novel splice forms, we compared the assemblies to a known annotation database, Ensembl\(^{15}\), using the program Cuffcompare from the Cufflinks package. This gave us between 35,000–52,000 transfrags per tissue that were associated with 13,000–17,000 Ensembl genes, of which a large fraction (between 5,000–20,000 per tissue, representing 11–45% of the total) appeared to be novel splice forms (Figure 1, Supplemental Figure S3 and Supplemental Table S3). Tissues with large numbers of new splice forms also had a larger fraction of candidate new splice forms.

Even with the best data and software, computational reconstruction of long transcripts from short reads is prone to assembly errors. We therefore focused on classes of alternative splicing events that are most likely to be assembled correctly. Exon skipping events are the most prevalent type of alternative splicing events in the human genome\(^{26}\), and are particularly easy to identify from transcript data and less likely to be mis-assembled. They have been extensively studied and are well represented in the databases. For these reasons, exon skipping events provide an excellent proxy for the number of other types of splicing variants in a sample.

We define an exon skipping ‘event’ as a pairing between an exon-containing form (‘on’) and an exon-excluding form (‘off’), occurring

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**Figure 1.** A high-level view of alternative splicing in sixteen human tissues: numbers of multi-exon ‘genes’ and transcripts from *de novo* transcript assemblies produced by Cufflinks (left), and by Cuffcompare (right). Since Cufflinks may break transcripts and genes into multiple fragments when there is insufficient read coverage, we used Cuffcompare to compare transfrags against the Ensembl reference annotations to produce a better estimate for the number of genes and transcripts in the samples. Results in the right panel show the total number of Ensembl annotated as well as novel genes, and respectively transcripts, found in each sample. The number of novel isoforms identified by Cuffcompare is shown in the bottom panel.
at the same exon and with the same flanking introns. The same exon (or intron) may be involved in multiple exon skipping events, though the number of such cases is small. To generate a catalog of events for the sixteen tissues, we analyzed transcript assemblies using our software ASProfile and identified differences in exon-intron structures characteristic of the various classes.

We found over 150,000 candidate alternative splicing events (Supplemental Table S4). Among these, we found 26,989 exon skipping events at 25,017 distinct exons, involving 22,145 distinct introns. Almost all of these events (25,920) were found in comparisons between different tissues, although a significant fraction (16,382) were also found when comparing isoforms within the same tissue. There were 1,069 instances of alternative splicing events that were restricted to a single tissue, most of them in testes (416) and brain (172).

Mapping artifacts can create false exon skipping events, due to incorrect or duplicated splice junctions or incorrectly reconstructed exons. To assess the accuracy of the data set and identify potential artifacts for future curation, we first looked for co-located events that showed small variations (≤5 bp) at exon and intron boundaries, which could be caused by imprecise mapping of spliced reads. Such variations could lead to redundancy in reporting the events. For reference, we compared the extent of variation against the ENSEMBL gene annotations. There were 1,822 (6.75%) events in our data set that represented slight variations of other events compared to 427 (1.7%) in the ENSEMBL data, suggesting that up to 5% of events in our data set may be redundant (Supplemental Figure S5 and Supplemental Table S5). However, this figure is likely an overestimate, given that small 5′ and 3′ exon splicing variations are hard to detect with conventional (Sanger) data and are likely underrepresented in the reference gene annotations. We also evaluated the reproducibility of our results when using other transcript assembly methods (IsoCERM, SLIDE, and Scripture), in a second test. We found that 84% (2,471 out of 2,934) of exon skipping events found in the adrenal sample alone were independently discovered when using one of the other transcript assembly methods (Supplemental Table S6). When aggregating data across the sixteen tissues, 92% (24,936) of the introns spanning skipped exons have at least two reads supporting them in the sixteen tissues; although in general exon-skipping introns have fewer supporting reads than other introns (Supplemental Figure S7). Similarly, in 21,469 (80%) of the exon skipping events, the exon was present in two or more tissues. Thus, while some assembly artifacts could still be present, most of the events discovered have strong supporting evidence.

How much alternative splicing is novel?

Simply counting the number of transcripts assembled from RNA-seq data is one way to measure the extent of alternative splicing. However, this can be confounded by transcripts that are assembled incompletely or incorrectly. Exon skipping events are discrete and easily counted, although it is worth noting that a given exon might be skipped in multiple distinct transcripts. To avoid the difficulties of counting all splice variants, we used the number of exon skipping events as a surrogate measure of splicing variation.

To identify which of the splice variants were previously unreported, we searched the 26,989 skipped exon events against four gene annotations databases: CCDS (23,353 sequences) (http://genom.ucsc.edu, download May 2011), UCSC Genes (73,671) (http://genome.ucsc.edu, download May 2011), Ensembl v.61 (120,122) (http://ensembl.org), and H-DBAS (58,609 mRNA and 37,096 fl-cDNA RASVs) (http://h-invitational.jp/h-dbas/, download May 2011). Importantly, these specific data sets and releases had been produced using almost exclusively traditional cDNA (EST, mRNA) resources, and therefore provide a fairly accurate assessment of the potential to discover novel alternative splicing variation in RNA-seq experiments. We found that over 60% (17,442) of the events were novel, even after allowing for slight differences in the annotation of exon boundaries present in the various databases (Supplemental Table S8). New exons, new introns, or both can lead to novel splicing events, but we discovered novel introns much more frequently than novel exons (2,914 exons and 15,958 introns). The majority of novel exons overlap known exons; i.e., one or both exon boundaries are novel, but not the entire exon. 884 exons did not overlap any previously annotated exon. A total of 996 (34.2%) of the novel exons and 3,801 (23.8%) of the novel introns were also supported by EST alignments, which provides independent cDNA evidence for those events.

One example of a novel event is shown in Figure 2. CHTOP (Chromatin target of Pritn1, synonym FOP) is a small nuclear protein on chromosome 1 characterized by an arginine- and glycine-rich region. It has a role in ligand-dependent activation repression of an estrogen receptor target gene32, and has been shown to be a critical modulator of gamma-globin gene expression33. The 84 bp in-frame exon at chromosome 1 positions 153,611,844–153,611,927, which we observed only in heart tissue, does not overlap any of the annotated structures for this gene and has only weak EST evidence, in the terminal exon of EST DB270513. However, the entire exonic region is highly conserved in placental mammals, strongly suggesting that this region is part of the spliced gene. Further, DNaseI hypersensitive sites lend support to an alternative transcript starting at this exon in thyroid tissue. This alternative transcription start site was also identified by our method, and is also missing from the annotation.

Another novel event occurs in the gene ASB15 (ankyrin repeat and SOCS box containing 15) (Supplemental Figure S9). Human ASB15 is known to be expressed predominantly in skeletal muscle and to participate in the regulation of protein turnover and muscle cell development by stimulating protein synthesis and regulating differentiation of muscle cells. Bovine ASB15 mRNA was also found in heart and pituitary gland tissue, and rat ASB15 was additionally present in kidney and lung tissue, but the amount in most other tissues analyzed was scarce34. These results are consistent with the Illumina Human Body Map data set. Here, exon chr7:123,257,633–123,257,718 is a novel shorter variant that shares its 5′ end with the annotated exon. Both the exon-containing and the exon skipping form are expressed in heart, and have strong read support in the 16 tissues (136 and 39 reads supporting the flanking introns, and 30 reads spanning the exon). Evidence for the novel splice junction is also present in skeletal muscle tissue. We
Figure 2. A novel alternatively spliced exon (chr1: 153,611,844–153,611,927) at the CHTOP gene locus, which does not overlap any known annotation. This novel 84-bp exon, marked with a red circle in the figure, is the 4th exon in one of the transcripts from heart tissue (heart.79763.3), and it appears exclusively in that tissue, although a partial form is present in a skeletal muscle transcript.
also found a novel putative intron retention event (chr7:123269489–123270019, 531 bp) whose sequence is conserved across multiple vertebrate species. Overall, our analyses underscore the vast potential for RNA-seq experiments to unearth novel splicing events and isoforms.

Characterization of exon skipping events

We next sought to characterize the set of exon skipping events within and across the sixteen tissues, which also offers a glimpse into the dynamics of alternative splicing in these tissues. We separately traced the presence of the two forms ('on' and 'off') to generate an alternative splicing profile for each tissue. For each event, we determined the exon inclusion ratio from the expression levels of isoforms containing the ‘on’ and the ‘off’ forms in that tissue: \( R = \frac{\text{FPKM}_{\text{on}}}{\text{FPKM}_{\text{on}} + \text{FPKM}_{\text{off}}} \), and then compared the profiles to determine similarities and changes in splicing patterns among tissues. We used the relative inclusion ratio to characterize such changes: \( \Delta = |R_i - R_j| \), and classified them based on the size of the difference. We separately trace exon skipping events that show large variation ('switches'; \( \Delta \geq 0.5 \)), essentially switching between a minor-form and a major-form, and those that show milder variation. Note that all of these evaluations, based as they are on a single sample from each tissue, provide only a qualitative assessment of variation. Multiple replicates would be required to make any conclusions about the statistical significance of these changes between tissues.

Of the 26,989 exon skipping events, between 10,000–20,000 are present in any given tissue. Most events (77–95%) have only one of the forms expressed in a given tissue, and only 5–23% have both forms present in the same tissue (Figure 3 and Supplemental Table S10). The exon-containing ('on') form is generally prevalent (\( R \geq 0.5 \)). When comparing the profiles between two tissues (Figure 4), 25–35% of the events show stable splicing patterns (\( \Delta < 0.1 \)), 5–10% are variable (0.1 \( \leq \Delta < 0.5 \)) and only 4–7% appear to switch. These proportions are quite similar among the tissues. Roughly 50–65% of the events are not comparable, with the event not found in either or both tissues. Further examination showed these to be due largely to the gene not being expressed (fragments per kilobase of transcript per million mapped (FPKM)<<0.1) or harboring different splice forms, whereas we expect the number of incomparable events caused by computational artifacts to be very low. A significant portion of these genes were expressed at low-to-medium levels (FPKM\leq10.0), which makes reconstruction difficult and may cause the event to be missed. (For an example, the comparison between the adrenal and adipose tissues is shown in Supplemental Figure S11). These observations suggest that both transcription and alternative splicing contribute significantly in shaping the transcriptomic differences among tissues, although more complete data sets and experiments are needed to be able to tease apart their specific contributions.

Characterization of novel events

We contrasted known and newly found events to determine characteristics that could have made the latter difficult to discover with conventional (Sanger) data and methods, and to derive insights into the types of experiments that can help fill in the gaps in the alternative splicing catalog.

First, we analyzed the variability in splicing patterns of events, distinguishing between 'switches' and events exhibiting milder variation. There was a slight but statistically significant difference between the distributions of novel and known events (chi-square 274.7; \( p = 0.0 \); Supplemental Table S12), with switches representing 69% of the known events and only 59% of the novel set.

Second, we assessed the tissue specificity of known and newly found exons and introns based on the data available (Figure 5). For this test, we binned both the known and the novel features according to the number of tissues in which they were found. Not surprisingly, novel exons and introns were significantly more likely to appear in a small number of tissues compared to their known counterparts, but the prevalence was remarkable for exons. For instance, while novel introns were more likely to belong to a single tissue by a 3.0:1 margin (48% versus 16%), that margin for exons was 5:1 (71% versus 14%). Considering that our search turned out many more novel introns than exons, this observation suggests that targeted studies will be needed in the future to identify these highly tissue-specific exons.

Discussion

Alternative splicing is a widely recognized RNA processing mechanism in eukaryotic species, playing a major role in the molecular biology of the cell, and within humans it has been implicated in multiple genetic disorders. The Human Genome Project created an initial map of splice variation more than a decade ago. However, despite concerted efforts over the following years, this map is still inaccurate and incomplete. The Ensembl annotation, which is among the most complete to date, currently contains seven variants on average per protein coding gene. This is likely an underestimate, as more variants are added every day. The challenge of cataloging all alternative splice variants is daunting, considering that every tissue and cell type can have a different transcriptome, further differentiated by the condition of the cell at the time it was surveyed.

Unlike traditional methods that have mined heterogeneous cDNA sequences collected over time, RNA-seq experiments can survey the transcriptome of a cell type or tissue at great depth, allowing characterization of alternative splicing in much finer detail than previously. The main drawback to RNA-seq today is that its shorter reads are more challenging to assemble into long isoforms. To avoid some of the uncertainty associated with transcriptome assembly, we focused here on alternative splicing events within a transcript, each of which can be detected with a single read.

We found over 150,000 candidate alternative splicing events, including roughly 27,000 exon skipping events, most of which (65%) were novel. New introns (15,958) were the main source of novel events in our data set, but we also found a large number of new exons (2,914). A large majority of the new exons appear to be tissue-specific, with 71% present in only one tissue, which may explain why they have not been detected previously. Tissue-specific exons represent a clear and important contribution of alternative splicing to tissue differentiation, hence it is noteworthy that 2,085 (38%) of the 5,520 events in our data set were newly identified in this study. Both novel exons and novel introns were more likely to be tissue-specific than those already in the public annotation sets. This suggests that targeted experiments in different tissues or cellular conditions will be more productive in identifying novel splice
forms in the future. This requirement is particularly relevant for identifying new exons, which have already been surveyed quite intensively, whereas even broad range RNA-seq experiments remain a rich source of new introns.

Our analysis of the 27,000 events across the sixteen tissues has also revealed insights into the dynamics of the alternative splicing repertoire and its role in tissue differentiation. With roughly 10–20% of the events showing variation across the tissues and 50–65% incomparable based on the existing data, the picture of alternative splicing contributions to tissue transcriptome differentiation vis-à-vis transcription is shaping up to be significant, albeit incomplete. Indeed, even in a deep and rich data set such as the Illumina Human Body Map, rare splice forms may be poorly represented or can be missed entirely. Also, our analyses here are based on a single experiment per tissue from a single individual, and therefore we cannot rule out polymorphic variation, although we expect its contribution to be small relative to tissue related differences\textsuperscript{2,14}. Of course, experimentally testing the events\textsuperscript{36} and

Figure 3. Splicing variation at skipped exon events as measured by the exon inclusion ratio $R = \frac{\text{FPKM}_{on}}{\text{FPKM}_{on} + \text{FPKM}_{off}}$ in the sixteen tissues. Most events within a given tissue are single variant (top). When both isoforms are present in a tissue, the exon is typically contained in the major form ($R \geq 0.5$) (bottom).
replication on multiple biological samples, from different individuals, will be essential for full validation.

While there are ongoing efforts to incorporate alternative splicing information from RNA-seq data into gene annotation databases, there is yet no repository specifically for human alternative splicing events. Our analyses have identified thousands of putative alternative splicing events, which we have compiled into a catalog of exon skipping events derived from RNA-seq data from multiple human tissues. This collection will be a valuable resource for investigating the mechanisms and evolution of alternative splicing, and as a complement to existing annotation databases. Although this catalog adds substantially to the list of known alternative splicing events, many more RNA-seq experiments will be needed to fully characterize alternative splicing over the full spectrum of tissue types and cellular conditions. Our methods, as implemented in the ASprofile software, are freely available to allow others to create similar databases for other organisms or experimental systems.

**Materials and methods**

**Sequence data**

RNA-seq data for the Illumina Human Body Map Project were downloaded from http://www.ebi.ac.uk/arrayexpress/browse.html?keywords=E-MTAB-513&expandefoo=on. For sequencing, samples for each of the 16 tissues (adrenal, adipose, brain, breast, colon, heart, kidney, liver, lung, lymph, ovary, prostate, skeletal muscle, testes, thyroid and white blood cells) were prepared by Illumina using their mRNA-Seq kit (Part #RS-100-0801). In brief, PolyA+ RNA was purified from 100 ng of total RNA with oligo-dT beads, and then fragmented with divalent cations under elevated temperature. First strand synthesis was performed with random hexamer and reverse transcriptase, and second strand synthesis with RNAseH and DNA PolI. Following cDNA synthesis, the double stranded products were end-repaired, a single “A” was added and then the Illumina PE adaptors were ligated on to the cDNA products. The ligation products were purified using gel electrophoresis. The target size range for these libraries was ~300 bp, such that the final library for sequencing would have cDNA inserts with sizes of ~200 bp long. One run of 2x50 bp paired-end sequencing was performed on the HiSeq2000 instrument, using one lane per tissue, to produce approximately 80 million read pairs per tissue (160 million sequences) (http://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-513/protocols/).

**Reconstructing the tissues’ transcriptomes**

To determine splice variants within each tissue, we aligned reads to the hg19 genome using TopHat v1.3.3 (parameters ‘-a 6 –F 0.05 –splice_mismatches=1 –max-multihits=10’). To allow TopHat to detect as many splice junctions as possible, we provided an intron database extracted from the UCSC known Genes data set (http://genome.ucsc.edu). Aligned reads were then assembled into transcript fragments using Cufflinks v0.9.3 (parameters ‘-F 0.05’). We used Cuffcompare to compare these transfrag to the Ensembl v61 annotation, and then Cuffdiff to redistribute reads along a high-confidence
set of transcripts obtained after eliminating likely artifacts and assemblies not associated with Ensembl genes. Cuffcompare classifies assembled transcripts into multiple categories in relation to reference transcripts, including equal, contained, new splice isoform, intron-located, pre-mRNA fragment, repeat, etc. We retained only transcripts that were deemed ‘equal’, ‘contained’, or ‘new splice isoforms’ as part of our high-confidence set for each tissue. FPKM expression level values for this set were then re-estimated from the original alignments using Cuffdiff.

Discovery of alternative splicing events

To determine alternative splicing events, we developed a software package, ASprofile, to analyze all pairs of transcripts in the sixteen tissues to determine exons included in one transcript and skipped in the other. We restricted the analysis to Ensembl genes across samples, which can be used to observe trends in the dynamics of alternative splicing profiles or to select promising candidates for laboratory testing. The software package is written in C and Perl and is available free of charge from the Zenodo repository (http://zenodo.org/record/7068; doi:10.5281/zenodo.7068) and from our web site at http://ccb.jhu.edu/software/ASprofile.

Comparison of alternative splicing events across tissues

For each event, we calculated the exon inclusion ratio \( R = \frac{\text{FPKM}_{\text{on}}}{\text{FPKM}_{\text{on}} + \text{FPKM}_{\text{off}}} \) for each tissue, similarly to Wang et al., where FPKM_{on} is the combined FPKM of all isoforms containing the ‘on’ form, and similarly for FPKM_{off}.”

Implementation

We implemented the methods in a software package, ASprofile, for discovering alternative splicing events in transcripts predicted from RNA-seq data and then comparing them across multiple conditions. ASprofile consists of programs for extracting (‘extract-as’), quantifying (‘extract-as-fpkm’) and comparing (‘collect-fpkm’) alternative splicing events. ‘Extract-as’ takes as input a GTF transcript file, for instance one produced by a transcript assembly program or a set of gene annotations, and compares all pairs of transcripts within a gene to determine exon-intron structure differences that indicate an AS event. The following classes of events are currently implemented: exon skipping, cassette exons, alternative transcript start and termination, retention of single or multiple introns, and alternative exon ends (Supplemental Figure S13). To determine alternative splicing events among multiple samples, a single input file must be created by concatenating the transcript files of individual samples, with the gene names a priori reconciled across the samples (for instance, by using the program Cuffcompare from the Cufflinks suite). The second program, ‘extract-as-fpkm’, calculates the FPKM of each event from those of transcripts harboring the event in a given sample, allowing for small variations (up to V bp, where V is a user-specified value) at the boundaries of the exons and introns. Lastly, the script ‘collect-fpkm’ collects the FPKM event values for all RNA-seq samples, and calculates and compares splicing ratios across samples, which can be used to observe trends in the dynamics of alternative splicing profiles or to select promising candidates for laboratory testing. The software package is written in C and Perl and is available free of charge from the Zenodo repository (http://zenodo.org/record/7068; doi:10.5281/zenodo.7068) and from our web site at http://ccb.jhu.edu/software/ASprofile.
Author contributions
LF and SLS conceived the project and designed the experiments. LF and LS carried out the computational analyses. LF wrote the first draft, and SLS edited the final version of the manuscript. All authors read and approved the final manuscript.

Competing interests
No competing interests were disclosed.

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Supplementary materials

Table S1. Total number of RNA-seq reads from each of 16 human tissues. “Mapped” refers to the number of reads that had at least one and no more than ten alignments to the genome.

| Tissue            | Reads   | Mapped  |
|-------------------|---------|---------|
| Adipose           | 154,600,144 | 138,872,338 |
| Adrenal           | 148,945,742  | 133,883,290 |
| Brain             | 147,026,094  | 134,791,012 |
| Breast            | 151,724,430  | 135,987,146 |
| Colon             | 164,874,886  | 150,966,150 |
| Heart             | 165,837,568  | 155,841,749 |
| Kidney            | 160,794,674  | 143,888,751 |
| Liver             | 160,097,246  | 149,817,522 |
| Lung              | 158,593,810  | 144,039,113 |
| Lymph             | 164,156,314  | 146,044,267 |
| Ovary             | 161,892,520  | 147,588,256 |
| Prostate          | 164,668,152  | 151,847,711 |
| Skeletal muscle   | 164,222,278  | 151,847,711 |
| Testes            | 163,672,398  | 149,727,704 |
| Thyroid           | 163,825,774  | 148,942,521 |
| White blood cell  | 162,434,296  | 149,167,286 |
| Total             | 2,557,366,326 | 2,333,252,527 |
Table S2. Overview of results from the Cufflinks transcript assembly process. Cufflinks assembles short reads aligned to a genome into a set of transcript fragments (‘transfrags’ or ‘transcripts’ or ‘txpts’, below), grouped by locus (‘gene’).

| Tissue   | Genes* | Multi-exon | Txpts/gene† | Transfrags | Single exon | Multiexon | Exons/txpt‡ |
|----------|--------|------------|-------------|------------|-------------|-----------|-------------|
| Adrenal  | 307,088| 23,135     | 2.0         | 357,774    | 311,766     | 46,008    | 5.6         |
| Adipose  | 134,348| 21,793     | 1.5         | 158,142    | 126,207     | 31,935    | 6           |
| Brain    | 239,971| 24,435     | 1.8         | 218,818    | 238,497     | 43,321    | 6           |
| Breast   | 191,620| 23,553     | 1.6         | 222,525    | 185,869     | 36,656    | 5.5         |
| Colon    | 116,723| 25,503     | 1.3         | 133,953    | 103,071     | 30,882    | 5.1         |
| Heart    | 149,951| 21,537     | 1.5         | 177,059    | 144,829     | 32,230    | 6.5         |
| Kidney   | 171,862| 24,000     | 1.5         | 197,742    | 162,910     | 34,832    | 5.4         |
| Liver    | 95,495 | 21,228     | 1.3         | 110,116    | 82,732      | 27,384    | 5           |
| Lung     | 172,439| 25,176     | 1.4         | 194,640    | 159,123     | 35,517    | 4.9         |
| Lymph    | 188,709| 23,766     | 1.5         | 216,692    | 179,938     | 36,754    | 4.9         |
| Ovary    | 210,035| 22,949     | 1.8         | 252,081    | 210,391     | 41,690    | 6.5         |
| Prostate | 161,526| 24,584     | 1.4         | 186,159    | 150,732     | 35,427    | 5.5         |
| Skel_muscle | 95,684 | 19,695 | 1.2        | 105,209    | 82,359      | 22,850    | 4.7         |
| Testes   | 220,459| 29,546     | 1.6         | 259,043    | 212,894     | 46,169    | 5.9         |
| Thyroid  | 187,169| 23,357     | 1.8         | 224,647    | 184,751     | 39,896    | 6.3         |
| White_blood | 98,731 | 19,943 | 1.3        | 113,656    | 88,538      | 25,118    | 5.5         |

*Genes containing multi-exon transcripts. †Averaged over multi-exon genes. ‡Averaged over multi-exon transcripts.

Table S3. Novel and known gene splice forms in the sixteen tissues, determined by comparison to the ENSEMBL v.61 annotation database using the program Cuffcompare. Cuffcompare compares each predicted transcript’s intron chain against those of the reference transcripts and classifies the transfrag as: ‘equal’ to a reference transcript, if their intron chains are identical; ‘contained’, if included in a reference transcript’s; or as a ‘new splice form’, if the intron chain has at least one splice junction in common with the reference transcripts (n.b., other Cuffcompare codes are not relevant and were omitted).

| Set       | Full-length (Equal) | Partial (Contained) | New splice forms |
|-----------|---------------------|---------------------|------------------|
| Adrenal   | 5,705               | 22,613              | 22,592           |
| Adipose   | 5,593               | 22,293              | 11,233           |
| Brain     | 6,497               | 20,673              | 19,329           |
| Breast    | 5,665               | 24,190              | 13,665           |
| Colon     | 4,689               | 28,715              | 9,310            |
| Heart     | 6,103               | 17,614              | 11,701           |
| Kidney    | 5,501               | 26,615              | 11,775           |
| Liver     | 4,483               | 24,563              | 7,692            |
| Lung      | 4,958               | 33,735              | 12,346           |
| Lymph     | 4,574               | 33,999              | 14,364           |
| Ovary     | 7,017               | 18,667              | 18,048           |
| Prostate  | 5,673               | 27,183              | 12,470           |
| Skel_muscle | 3,318 | 33,999 | 4,999 |
| Testes    | 7,077               | 23,648              | 17,306           |
| Thyroid   | 6,487               | 21,098              | 16,667           |
| White_blood | 4,568 | 25,004 | 6,533 |
Figure S3. Proportion of predicted novel splice forms from the total number of transcript assemblies, including novel and known (top), and absolute counts of novel splice forms (bottom) detected in the 16 tissues. Novel and known isoforms are determined by comparison with ENSEMBL v.61 gene annotations using the program Cuffcompare (see caption to Table S3 for more details).

Table S4. Numbers of candidate alternative splicing events found in the Illumina Human Body Map data, by type.

| Event type                        | Code | Count |
|-----------------------------------|------|-------|
| Exon skipping (single exon)       | SKIP | 26,989|
| Exon skipping (multiple exons)    | MSKIP| 13,814|
| Alternative 5’ terminal exon*     | TSS  | 97,399|
| Alternative 3’ terminal exon*     | TTS  | 45,609|
| Intron retention (single intron)  | IR   | 16,671|
| Intron retention (multiple introns)| MIR  | 2,636 |
| Alternative exon boundaries       | AE   | 22,767|

*Includes 17,200 ‘default’ TSS (TTS) sites for 17,200 genes.

S5. Potential effects of imprecise splice junctions on the exon skipping (ES) estimates

One class of mapping artifacts that can contribute to creating false exon skipping events is that of ‘imprecise splice junctions’, where the same intron in different spliced reads is mapped by the alignment software at slightly different positions, typically within a few bp of each other, thus creating the appearance of several introns at that location. Such variations could lead to redundancies in reporting exon skipping events. To estimate the potential effects of imprecise splice junctions on our data set, we searched the 26,989 events against each other to identify those that are co-located and differ only slightly (by some maximum number of bases, \( V \) bp) in their exon-intron boundaries. (Note, however, that such events would also include all true alternative 5’ and 3’ exon ends). At each locus, we designate one event as ‘anchor’ (true), and refer to the co-located events as ‘variations’. We repeated the operation for 25,114 events discovered from the Ensembl v. 61 annotation with our software ASprofile, to create a baseline for comparison. For our data set of 26,989 events, labeled ‘Cufflinks’ in Figure S5 below, small \( \leq 5 \) bp differences in splice junctions account for the majority of ES variations, beyond which their value as a fraction of the total number of ES events becomes relatively stable. In contrast, the percentage of Ensembl v. 61 event variations continues to increase roughly linearly with the distance from the ‘anchor’ splice junction. Comparing the two curves, we estimate that up to \(~1,400\) of ES events (5%) in our data set could be redundant (column 10 in the Table S5). We expect this value to be an overestimate, given that true small 5’ and 3’ exon alternative splicing variations are poorly represented in annotation databases.

S6. Reproducibility of exon skipping event discovery with other assembly methods

To assess the reproducibility of our event discovery method, we compared the exon skipping events discovered from transcripts assembled with Cufflinks and with three other reference transcript assembly algorithms (Scripture\(^29\), IsoCEM\(^27\) and SLIDE\(^28\)). We applied each program to the adrenal sample of the Human Body Map data and used our program ASprofile to extract exon skipping events from the assembled transcripts.

As the results in Table S6 indicate, when compared to Cufflinks, IsoCEM and SLIDE find roughly half the number of events, whereas Scripture finds roughly 60% more events. (Allowing for small differences, up to \( V \) bp, in the coordinates of introns and exons to account for imprecise mapping of splice junctions, has negligible effects on the comparison results). There is generally good agreement among the methods, as most of the events predicted with one method can also be retrieved by at least one of the other methods. In particular, 84% (2,471 out of 2,934) of the events predicted when using Cufflinks can be confirmed by other methods. Additionally, our analysis reveals a number of potentially new events, most of them discovered by Scripture, which could be used to enrich our repository in the future. However, each program has its own characteristics and biases, and a careful analysis has to be performed before incorporating these events into the database.
Table S5. Comparison of co-located ES events among the 26,989 events discovered from the Illumina Human Body Map data and the 25,114 exon skipping (ES) events extracted from the Ensembl v.61 annotations. All – all events co-located within V bp; Loci – clusters of co-located events (each with a representative ‘anchor’); Var (=All-Loci) – variations (not including ‘anchors’), representing potential mapping artifacts.

| V     | Cufflinks (26,989 events) | Ensembl 61 (25,114 events) | % Diff | Potential artifacts |
|-------|---------------------------|----------------------------|--------|---------------------|
|       | All | Loci | Var. | %   | All | Loci | Var. | %   |               |
| 1     | 647 | 300  | 347  | 1.29 | 90  | 45   | 45   | 0.18 | 1.10 | 298          |
| 2     | 1271| 585  | 686  | 2.54 | 143 | 71   | 72   | 0.29 | 2.26 | 608          |
| 3     | 2394| 1153 | 1241 | 4.60 | 530 | 262  | 268  | 1.07 | 3.53 | 953          |
| 4     | 3079| 1476 | 1603 | 5.94 | 724 | 359  | 365  | 1.45 | 4.49 | 1210         |
| 5     | 3507| 1685 | 1822 | 6.75 | 839 | 412  | 427  | 1.70 | 5.05 | 1363         |
| 6     | 3803| 1832 | 1971 | 7.30 | 894 | 440  | 454  | 1.81 | 5.49 | 1483         |
| 7     | 3923| 1890 | 2033 | 7.53 | 930 | 456  | 474  | 1.89 | 5.65 | 1523         |
| 8     | 4003| 1929 | 2074 | 7.68 | 959 | 471  | 488  | 1.94 | 5.74 | 1549         |
| 9     | 4126| 1991 | 2135 | 7.91 | 1020| 502  | 518  | 2.06 | 5.85 | 1578         |
| 10    | 4192| 2057 | 2135 | 7.91 | 1068| 500  | 568  | 2.26 | 5.65 | 1524         |

Figure S5. Number of event variations due to small differences in exon and intron boundaries as a fraction of the total number of events, with varying cutoffs for the allowable difference (V bp). Values shown are for exon skipping (ES) events derived from the Illumina Human Body Map data (“Cufflinks”) and from Ensembl v.61 gene annotations, from columns 4 and 8 in Table S5.

Table S6. Reproducibility of exon skipping (ES) discovery from transcripts assembled with different methods, and robustness with varying cutoffs (V bp) for the margin of error when comparing exon and intron boundaries.

| Method  | ES events | Found by other methods |
|---------|-----------|------------------------|
|         | V=0      | V=5       | V=10      |
| Cufflinks| 2,934    | 2,471     | 2,480     | 2,482    |
| IsoCEM  | 1,486    | 1,361     | 1,364     | 1,365    |
| SLIDE   | 1,418    | 860       | 884       | 895      |
| Scripture| 4,770    | 2,984     | 3,072     | 3,096    |
Figure S7. Histograms of reads in the 16 tissue samples supporting the two introns flanking the alternatively spliced exon (blue, red) and the intron spanning the exon (green), respectively, for the 26,989 identified exon skipping events. (A) Diagram of an exon skipping event, illustrating the three types of introns: left – blue, right – red, and intron spanning – green. (B) Read histograms for the three categories of introns. The x-axis represents the number of supporting reads for an intron, grouped in bins, and the y-axis shows the percentage of introns by levels of supporting reads (in bins).

Table S8. Summary of novel features by comparison to four annotation databases (CCDS, UCSC Genes, Ensembl v. 61, and H-DBAS). Exon and intron boundaries were compared allowing for a small difference (up to V=0, 5 and 10 bp). Values in bold are those reported in the main text.

| Novel feature  | V=0 | V=5 | V=10 |
|----------------|-----|-----|------|
| Exon+Intron    | 2,518 | 1,430 | 1,214 |
| Exon only      | 1,954 | 1,484 | 1,368 |
| Intron only    | 13,531 | 14,528 | 14,726 |
| Known          | 8,986 | 9,547 | 9,681 |
**Figure S9. Novel exon skipping event at the human ASB15 gene locus.** Exon chr7:123,257,633–123,257,718, present in heart (heart.653543.1 and heart.653543.2) and with partial support in skeletal muscle tissue (skel_muscle.206929.2), is novel. Red arrows point to the exon appearing in two isoforms reconstructed in the heart sample, and to the (potentially) partial form in a skeletal muscle transcript for the ASB15 gene. The spanning intron in the heart sample is also novel. Additionally, we found a potentially retained intron (chr7:123,269,489–123,270,019; thyroid.843995.1), circled in red, whose 531 bp sequence is conserved across multiple vertebrates.
Table S10. Exon skipping events expressed in each tissue sample, and number/percentage of these events that have only one expressed isoform in the sample.

| Tissue            | Events | Single variant |
|-------------------|--------|----------------|
| Adrenal           | 16,736 | 13,403 (80.1%) |
| Adipose           | 16,045 | 13,819 (86.1%) |
| Brain             | 17,415 | 13,865 (79.6%) |
| Breast            | 16,087 | 13,870 (86.2%) |
| Colon             | 13,802 | 12,464 (90.3%) |
| Heart             | 17,048 | 14,769 (86.6%) |
| Kidney            | 15,319 | 13,785 (90.0%) |
| Liver             | 12,390 | 11,388 (91.9%) |
| Lung              | 13,736 | 12,310 (90.6%) |
| Lymph             | 13,393 | 11,765 (87.8%) |
| Ovary             | 19,209 | 15,052 (78.4%) |
| Prostate          | 16,011 | 14,316 (89.4%) |
| Skel_muscle       | 10,160 | 9,614 (94.6%)  |
| Testes            | 19,124 | 15,127 (79.1%) |
| Thyroid           | 18,689 | 14,362 (76.8%) |
| White_blood       | 12,954 | 12,017 (92.8%) |

Figure S11. Expression levels of genes for the 13,946 events not comparable between adrenal and adipose tissues. 2,085 (15.0%) event genes are not expressed (FPKM<0.1), whereas 3,487 (33.7%) have FPKM values less than 10.0 and therefore may be incompletely reconstructed, which can cause a splice form to be missed.
Table S12. Variability of novel exon skipping events compared to known events. Novel events are depleted in ‘switches’ compared to known events (test of homogeneity, df=2, chi-square 274.7, p = 0.0).

| Events | Stable | Variable | Switches | Total |
|--------|--------|----------|----------|-------|
| Novel  | 1,788  | 5,334    | 10,320   | 17,442|
| Known  | 688    | 2,241    | 6,618    | 9,547 |
| Total  | 2,476  | 7,575    | 16,938   | 26,989|

Figure S13. Classes of alternative splicing events detected by ASprofile by pairwise transcript comparisons. (A) Exon skipping (SKIP) and cassette exons (MSKIP); (B) retention of single (IR) and multiple (MIR) introns; (C) alternative exon ends (AE); (D) alternative transcription start site (TSS); and (E) alternative transcription termination site (TTS). Alternatively spliced features are shown in red.

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8. Gonzalez-Porta M, Calvo M, Sammeth M, et al.: Estimation of alternative splicing variability in human populations. Genome Res. 2012; 22(8): 1228–1238. PubMed Abstract | Publisher Full Text | Free Full Text
In this article, the authors address the question of how much alternative splicing can be identified in 16 different human tissues, and how much of the alternative splicing is specific for one or another tissue. To do so, they analyzed 50-bp paired-end sequences from 16 different tissue samples sequenced relatively deeply (ca. 80 million read pairs per tissue sample). The data has been made available by Illumina as the Human Body Map project. Computational analysis was then performed with the programs TopHat and Cufflinks. This type of analysis assembles the reads into a set of transcripts that are compatible with the splicing patterns inferred from reads that are split over multiple exons. Between 23,000–46,000 partial or complete transcript assemblies (“transfrags”) were obtained per tissue. Approximately 5,000–20,000 transfrags per tissue, or 11–45% of the total, were identified as potentially novel splice forms. As the authors note, it is still difficult to assemble complete transcripts from relatively short RNA-seq reads in this way, and they thus make the very reasonable decision to concentrate their further analysis on alternative splicing events that are likely to be valid. Thus, it is easier to analyze the spectrum of exon skipping/inclusion events at a particular exon.

In general, the article is well done and will be of interest to those involved with RNA-seq or alternative splicing. One particularly interesting, if not entirely surprising, finding of this study is the fact that the majority of the novel exon insertion events showed a high degree of tissue specificity, thus providing an obvious explanation of why the corresponding exons are not yet in public databases. This suggests that efforts should be spent to characterize the full range of splicing in tissues in order to understand their biology.

Comments:

1. It would be useful if the authors could provide more discussion of the current state of the art of transcript identification using RNA-seq reads. In particular, the authors discuss three other transcript assembly programs in the supplement (S6), finding that they identify 50-60% of the isoforms called by Cufflinks. Have the authors performed their downstream analysis using the results of one of these programs? How different would the results be?

2. The authors should try to place their results in the context of many other efforts at identifying novel isoforms from the literature and discuss the relative merits of their approach.

3. It would be useful to have a table with quality metrics for this dataset, e.g., Phred scores, percent of mapable reads etc. - table S1 could be extended for this purpose.
4. Although much of the results deal with exon skipping/insertion events, it would be interesting to
hear a little about results for other classes of alternative splicing that can be identified by the
authors’ software, e.g. alternative transcription start and termination sites.

5. In the results section, it would help the presentation of the material to provide a brief description of
the relative inclusion ratio at this portion of the text and to define what i and j are (i.e., take some of
the material from the methods section that explains this).

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

Competing Interests: No competing interests were disclosed.

Author Response 18 Nov 2013

Liliana Florea, Center for Computational Biology, McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University, USA

We thank Peter Robinson for his insightful comments and suggestions, which we address in the following:

- Transcriptome assembly represents a complex topic on its own, and is only a tangent to our
work, therefore we believe that it would be best addressed in detail elsewhere. We wish to
point the reviewer and the readers to a review on current transcript assembly methods that
we wrote recently, to appear in IEEE/ACM Transactions in Bioinformatics and
Computational Biology journal. We also included a brief evaluation of several transcript
assemblers on a simulated data set (new Table S2), showing that Cufflinks is the only
program among those tested that can identify alternative splicing events with high enough
accuracy (~90%) to allow meaningful downstream analyses, though we will explore
combinations of these programs in the future. Minor correction to the statistic quoted by the
reviewer: 84% of the Cufflinks-predicted exon skipping events are reproducible by other
methods.

- There is indeed a rich body of work in the area of alternative splicing. Notably, two early
studies (references 1 and 2) highlighted the extent of alternative splicing in the human
genome based on analyses of RNA-seq data, albeit those data sets were not nearly large
enough to be able to characterize it in detail, and a more recent study (Merkin et al., 2012)
extended those analyses in the context of multiple species. Similarly, there are now several
methods (e.g., MISO, SpliceTrap and MATS) that infer or quantify exon skipping events
directly from RNA-seq reads, and therefore are complementary to our approach. Using a
very large data set, our work provides a more detailed picture of alternative splicing variation
and produces both an easy-to-use tool and a database that we believe will be valuable in
studies of alternative splicing mechanism and function.

- We added percentage of mapped reads and number of properly paired reads to Table S1,
as indicators of read alignment quality.

- We think that analyzing other types of events is indeed a very interesting experiment, which
due to scale we plan to address in our future work.
The definitions were already in the main text (page 7, section “Characterization of exon skipping events”), but we have clarified i and j.

**Competing Interests:** None

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**Manuel Corpas**
The Genome Analysis Center, Norwich, UK

The article 'Thousands of exon skipping events differentiate among splicing patterns in sixteen human tissues' presents a systematic analysis of alternative splicing of the Illumina Body Map RNA-seq set, probably the most complete set of RNA-seq data to date. This data set consists of 160M reads each for 16 tissues, each from a different individual.

From the standpoint of a reviewer I consider this article to be of very good quality, and have very much enjoyed reviewing it. There are a few suggestions I hope could help improve this work.

1. P3. pp.2: Has alternative splicing profiling been attempted before? If so, could this please be shown in the background section? If not, how does this technique specifically differ from what has been done before? Some more context would be appreciated.

2. P3. pp.3: What is the effect of comparing tissues from 16 different individuals? I would expect that the genetic variation of each individual would have some effects in the actual AS patterns observed when compared. I see that these samples belong to people with different ethnic backgrounds and ages. I would be interested to know what the authors think about how different results would be if all 16 tissues had been from one individual.

3. I tried to compile ASprofile on a mac and I got this error:

   libc.h:15:20: error: malloc.h: No such file or directory

   I solved this by changing this line with #include <stdlib.h>. Perhaps you might want to add these lines, so that mac users do not encounter the problem:

   #if !defined(__APPLE__)
   #include <malloc.h>
   #endif

4. Although the license for ASprofile is available in the Zenodo page, it might also be a good idea to mention it somewhere in the article. Not mentioning a license may make some users uneasy, even if it says it is open source.

5. In S1, how many of the mapped reads were properly paired?
6. Why did you use such old versions of Ensembl genes (Ensembl 61, February 2011), UCSC Genes, CCDS genes and H-DBAs? I also find that the TopHat (v1.3.3) and Cufflinks (v0.9.3) versions are quite old. The current version for TopHat is 2.0.9 and the current version for Cufflinks is 2.1.1. The current version of Ensembl is version 73, September 2013. The usage of such old versions concerns me a little, particularly those of the gene annotations, given that a lot of the results that are presented in this work rely heavily on the comparison of AS profiles against those annotations. Some of the results might be different if a newer version of the annotations were up-to-date.

7. P5, pp.5: The example of a novel event in CHTOP does not provide clear evidence as to how many reads support the novel exon. Please provide some numbers.

8. P7, pp1: Please could you provide evidence for the novel putative intron retention event located in the chr7:123269489-123270019 region?

9. In Figure S7.B, I would appreciate some text describing the biological significance of having the 100 bin in the x-axis with the greatest bars. Overall I am unable to understand what this figure means.

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

**Competing Interests:** No competing interests were disclosed.

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**Author Response 18 Nov 2013**

Liliana Florea, Center for Computational Biology, McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University, USA

We wish to thank Manuel Corpas for his thorough review and suggestions. In responding to his comments:

- We would like to point to our answer to question 2 from Reviewer 1, where we more broadly address this and other related questions.

- An earlier study by Wang *et al.* (reference 2) also tackled polymorphic splicing differences, finding that these represent a much smaller fraction of the AS variation compared to tissue differences. Therefore, while some polymorphic differences exist, they are not likely to distort the picture we paint in this study. We agree that analyzing data from multiple tissues in the same individual would be highly informative and we are actively looking for suitable data sets, which are not available at the moment.

- We thank the reviewer for taking the time to test the software. We updated the code and included a note on the license (GNU GPL) in the text (page 3).

- We have included proper read pairing information for all tissues in Table S1.

- With respect to the versions of the software, we note that at the time we ran our analyses we used the latest versions of each of these programs and databases, however our paper was caught up in the reviewing process at another journal for over a year. After two rounds of very slow reviews and multiple requests for revisions, the other journal was still not satisfied and therefore we decided to submit to F1000Research, which has far faster publication...
turnaround. We should emphasize that none of the previous reviewers’ comments questioned the validity of our results, and our revisions did not substantively change any of our conclusions. Nevertheless, we also searched the ‘novel’ events against the aggregate set of Ensembl 73, GENCODE v.17 and RefSeq representing the most recent versions of these collections, and only found 21 exons and 9 introns (note that these did not include the events depicted in Figures 2 and S9). Hence, the results discussed in the manuscript still hold.

- We have added read support information for the exon skipping event in Figure 2, and for the putative intron retention event in Figure S9.

- The ‘100’ bin in Figure S7 (now S8) shows the proportion of introns supported by more than 100 reads, for each of the three types of introns. Most events have deep support (>=100 reads) for the flanking introns, and to a lesser extent for the spanning intron, which is consistent with our earlier finding that the exon tends to be skipped in the minor form (Figure 3).

**Competing Interests:** None