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Improved Annotation of 3' Untranslated Regions and Complex Loci by Combination of Strand-Specific Direct RNA Sequencing, RNA-Seq and ESTs

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Abstract

The reference annotations made for a genome sequence provide the framework for all subsequent analyses of the genome. Correct and complete annotation in addition to the underlying genomic sequence is particularly important when interpreting the results of RNA-seq experiments where short sequence reads are mapped against the genome and assigned to genes according to the annotation. Inconsistencies in annotations between the reference and the experimental system can lead to incorrect interpretation of the effect on RNA expression of an experimental treatment or mutation in the system under study. Until recently, the genome-wide annotation of 3' untranslated regions received less attention than coding regions and the delineation of intron/exon boundaries. In this paper, data produced for samples in Human, Chicken and A. thaliana by the novel single-molecule, strand-specific, Direct RNA Sequencing technology from Helicos Biosciences which locates 3' polyadenylation sites to within +/- 2 nt, were combined with archival EST and RNA-Seq data. Nine examples are illustrated where this combination of data allowed: (1) gene and 3' UTR re-annotation (including extension of one 3' UTR by 5.9 kb); (2) disentangling of gene expression in complex regions; (3) clearer interpretation of small RNA expression and (4) identification of novel genes. While the specific examples displayed here may become obsolete as genome sequences and their annotations are refined, the principles laid out in this paper will be of general use both to those annotating genomes and those seeking to interpret existing publicly available annotations in the context of their own experimental data.

Introduction

There are two key features to a genome: the underlying sequence made up of the four nucleic acids (A, C, G, T), and its annotation. The majority of applications of a genome sequence rely on the gene structures and associated features provided by the reference genome annotation. Methods to annotate a newly sequenced genome are well developed and exploit both data-driven and ab initio feature prediction [1,2], but annotation is always derived from a snapshot of knowledge at the time it is carried out. As new data become available, the annotation must be revised if it is to remain relevant and useful (e.g. [3–6]). Annotation projects for the most complete and well described metazoan genomes: human[7]; mouse[8] and zebrafish[9], combine automatic methods with manual curation to provide an authoritative annotation that is regularly updated by incorporating new experimental data (e.g. [10]). The reference annotations for most other genomes rely more heavily on fully automatic annotation with limited manual curation. Since the structure of the gene transcript can vary according to cell type, treatment and other stimuli, the annotation that is most relevant may need to be re-defined for each set of experimental conditions. Advances in short-read, high-throughput transcript sequencing (RNA-seq) and its use in differential expression analysis have highlighted the importance of accurate gene models and prompted the development of methods to carry out experiment-specific predictions of gene structure (e.g. see [2,11–14]). However, conventional RNA-seq experiments often do not define the ends of genes with high precision. Incorrect assignment of the 5' and 3' UTRs may cause reads in an RNA-seq experiment to be assigned to intergenic regions and so give erroneous estimates of gene expression. Furthermore, the short read length may not provide evidence for
an unambiguous gene structure where there are overlapping genes, while RNAseq data that are not strand-specific are complex to apply in areas where genes overlap.

Recently, techniques have been developed that allow sites of cleavage and polyadenylation at the 3' end of transcripts to be identified in a high-throughput manner. These include 3P-Seq which has been applied to the characterisation of 3'UTRs in C.elegans [15] and zebrafish [16] and Helicos Bioscience's single-molecule direct RNA sequencing (DRS) [17] which has been applied to large-scale 3'UTR studies in human [18] A. thaliana [19], and yeasts [20,21]. DRS [17] captures RNA by the poly(A) tail and sequences the RNA immediately adjacent, so giving a very clear read-out of the transcript's 3'-end. DRS is strand-specific, has no amplification step, is less susceptible to internal priming than other methods and since it sequences RNA not DNA, does not require reverse transcription and the artefacts that can generate.

DRS has already been used in an automatic protocol to re-annotate the 3'-ends of over 10,000 protein coding genes in A. thaliana of which more than 3,400 were extended by at least 10 nt. [19]. Prior to the introduction of high-throughput sequencing technologies, expressed sequence tag (EST) libraries were commonly used to inform and validate gene models. Large libraries were produced since each tag only gave information about parts of a gene. Despite their size, EST libraries were often incomplete and error-prone due to PCR and reverse transcriptase artefacts. In contrast, RNA-seq datasets cover the vast majority of the transcriptome, but are based on shot-gun sequencing which requires reconstruction of the short reads. Typically, RNA-seq data does not retain strand information of the parent mRNA molecule.

In this study the potential of combining DRS with conventional RNA-seq, small RNA-seq (sRNA-seq) and archival expressed sequence tag (EST) data for genome annotation in human, chicken and A. thaliana is explored. Combining DRS, RNA-seq, EST and sRNA-seq data promises to mitigate the limitations of each individual technology; providing multiple, orthogonal, sources of evidence for gene intron/exon structure, 3' UTR regions and mature small RNAs and microRNAs, even in complex genomic regions.

**Materials and Methods**

In this paper, data from the authors' own laboratories were combined with data from public archives. The findings in this study are based on data produced from multiple collaborations and the choice of species reflects the data available rather than a specific design. The source of all data presented here is described below.

**Gallus gallus (chicken) DRS Data**

Sample Dissection. Pre Neural Tube (hereafter PNT) explants were dissected from Hamburger and Hamilton stage 10, 10 to 12 somite chick embryos ([22]). The explant was taken from a region rostral to the node and at a two presumptive somite distance from the last somite formed (somite I). The notochord was removed by controlled trypsin digestion aiming to keep the neural ventral midline. Dissections were carried out in L15 medium at 4°C and explants were taken for RNA extraction and DRS sequencing from three individual embryos (biological replicates).

RNA Extraction & Quality Testing. All surfaces and dissecting tools were treated with RNazap (Ambion) and rinsed with DEPC-treated water. RNA was extracted from the three PNT explants in Trizol reagent (Invitrogen) by phase separation with chloroform, followed by precipitation with isopropanol and linear acrylamide. The RNA was washed in 70% ethanol, air-dried, re-suspended in DEPC-water and frozen in liquid nitrogen.

Total RNA was quantified and quality tested using the Agilent RNA assay (Agilent Bioanalyzer pico RNA chip) by Helicos Biosciences. Samples with a RIN number above 8.0 were selected, and were then sequenced by DRS ([17]); producing 7.2-16.4 million raw reads per sample.

**DRS Data Processing.** Raw DRS reads from each sample were mapped to v2.1 of the chicken genome (Galgal3) with Helicos Biosciences' open-source mapping pipeline Helisphere (v2.0.022410) with the default parameters. The mapped reads were then filtered with four additional selection criteria to remove as much noise from the data as possible. Only reads with unique, high-quality, mappings to the genome (both locally and globally) were accepted. DRS sequencing technology is prone to producing reads that require a large number of insertions or deletions (in-dels) to align to the genome ([17,23]). Accordingly, to minimise ambiguity, only reads whose best-match alignments contained fewer than four indels, and whose read length was greater than 25 bases were accepted. Finally, all reads that map to any positions in the genome with fewer than 3 reads coverage per replicate were discarded. This threshold was chosen close to require at least three reads at a given genomic position in each of three replicates, ensuring that the retained peaks were reproducible across the three replicates and that they had a total signal-to-noise ratio (based on Poisson counting statistics) of < = 3.

Based on the existing chicken genome annotations from Ensembl, this resulted in a total of ~5,178 Ensembl genes with measured expression in all three PNT DRS replicate datasets. Data are available from www.compbio.dundee.ac.uk/polyadb and will be deposited at the European Nucleotide Archive.

**Gallus gallus Illumina RNA-seq Data**

The publicly available chicken Illumina RNA-seq data discussed here forms part of a study that examined gene expression in mammalian organs (Short Read Archive study: SRP007412 GSE30352 - [24]). This study used the Illumina Genome Analyser IIx platform to generate 76 bp reads for six tissues (brain - cerebral cortex or whole brain without cerebellum, cerebellum, heart, kidney, liver and testis) from one male and one female per somatic tissue (two males for testis). Data for the chicken were generated for this mammalian-focussed study as an evolutionary outgroup. The data were downloaded from the Short Read Archive, converted to fastq format with the SRA toolkit (v2.1.10, http://www.ncbi.nlm.nih.gov/Traces/sra/sra.cgi?view = software). The reads in each dataset were then aligned to v2.1 of the chicken genome (Galgal3) with the splice-aware alignment software TopHat (v2.0.0, http://tophat.cbcb.umd.edu/ - [14]) in conjunction with Bowtie (v2.0.0 beta3, http://bowtie-bio.sourceforge.net/index.shtml - [13]), with the --coverage-search, --micronox-search and --b2-very-sensitive options in addition to the TopHat defaults. Combined, the twelve samples total ~251 M reads, 64% (~161 M reads) of which map to the genome using these settings. Remapping to Galgal4 raised the total proportion of mapped reads to 69% but did not significantly affect the annotation examples shown in this paper.

**Homo sapiens skin DRS data**

Sample Dissection. A clinically normal human skin sample was obtained by 4 mm punch biopsy of skin tissue removed during plastic surgical procedures from the abdomen of an adult female, with approval from the local Research Ethics Committee, under the governance of Tayside Tissue Bank. The biopsy sample was
non-unique alignment hits were discarded and one hit selected at rand if there were several non-unique local hits found in a genetic region. Reads with more than four indels were discarded and read alignments refined by an iterative multiple alignment procedure while DRS reads containing low complexity genomic regions, as identified by DustMasker from the Blast+ 2.2.24 package, were discarded, as previously described [19]. These additional filters reduced the fraction of potentially incorrect alignments. The data have been deposited European Nucleotide Archive (ENA): Study, PRJEB3993; accession no, ERP009245.

Arabidopsis thaliana RNA-seq data

RNA-seq reads available in the accession SRR394092 were taken from the European Nucleotide Archive. These reads were generated from total RNA extracted from 10 day-old seedlings of A. thaliana (Columbia-0 ecotype) and sequenced by Illumina HiSeq 2000. All details of material preparation are described [27]. The 51.8 M raw reads length of 50 bp were aligned with the splice-aware alignment software TopHat v2.0.0 (this version of TopHat uses Bowtie v2.0.0 beta5) with the —b2-very-sensitive option in addition to the TopHat default options against the TAIR10 release of the A. thaliana genome. The total number of uniquely aligned reads was 48.8 M (94.2% of the raw reads).

Arabidopsis thaliana small RNA-seq data

Publicly available small RNA-seq data were taken from the European Nucleotide Archive (accession number is SRR167709). Total RNA for these data was extracted from immature flowers of wild-type A. thaliana (Columbia-0 ecotype), processed with Illumina Small RNA Sample Prep Kit and sequenced with HiSeq 2000. All details of material preparation are described [27]. The accession consists of 34.2 M of 36 bp non-aligned reads. The raw reads were quality-clipped, had their adapter sequences removed and remaining reads shorter than 16 bp were discarded as previously described [26]. The remaining 12.7 M reads were collated as 6 M unique sequences for alignment to the genome. The sequences were aligned to the TAIR10 release of the A. thaliana genome with bowtie v0.12.3 (parameters: -a —best —strata -v f). Bowtie was chosen, rather than bowtie2, as it is more suitable for small RNA-seq where gaps are of less relevance and reads are <50 bp in length.

Arabidopsis thaliana EST data

The A. thaliana EST data available in IGB were taken from the PlantGDB resource which aggregates the EST sequences from GenBank’s nucleotide database and splits them by species. The sequences used here are from GenBank version 187. They can be downloaded in fasta format from ftp://ftp.plantgdb.org/download/FASTA_187/EST/Arabidopsis_thaliana.mRNA.EST.fasta

Results

In this work, the definitions of ‘gene’ and ‘gene-associated regions’ (GARs) as suggested by Gerstein and colleagues [29] are followed. The results are divided into four sections where the major strengths of combining DRS data with other high-throughput transcriptomics data are highlighted by nine examples of feature re-annotation of genes and their GARs. Section 1 focusses on how the broad-coverage of RNA-seq and EST data help to bridge the gap between existing annotations and the DRS read data, enabling improved annotation of transcribed, polyadenylated regions. Section 2 illustrates how the positional specificity and native stranded-ness of DRS data enable re-annotation of complex genomic regions, without which the RNA-seq data could
not be used effectively either for re-annotation or further downstream analysis. Section 3 examines the synergy between standard RNA-seq, DRS and sRNA-seq data in providing a more complete picture of non-coding RNA expression than any of these datasets can provide individually. Section 4 briefly considers the potential for combined data to enable the discovery of new genes.

Section 1: Gene and 3' UTR re-annotation by combining DRS and RNA-seq data

A simple example: Chicken BMPR1A. The chicken genome sequence and gene models based on EST data were first released in 2004 (International Chicken Polymorphism Map[28]) with a second, more complete revision (v2.1) released 2006. A draft update to v2.1 was released in 2012, but this is yet to be annotated fully. Accordingly, most current research relies on v2.1 and its annotations and does not take account of evidence from DRS experiments.

Figure 1 shows the genomic context and information sources around BMPR1A, a gene important in development (FPI5350_CHICK, ENSGALG0000002905; [30–32]). The annotation of this gene and its GARS differ between Ensembl and RefSeq. Ensembl presents a single gene model and two short novel protein coding models. The canonical transcript (ENSGALG000000040319, see Table 1) covers 39,340 bp with twelve exons of 100298 bp, and an associated 228 bp 3' UTR. In contrast, the RefSeq annotation covers 39,340 bp, including a 21 bp longer first exon and a 17 bp shorter 3' UTR. Although the basic gene intron/exon structure and the 5' UTR are annotated in Ensembl/RefSeq, no 3' UTR is present in the RefSeq annotation and the 3' UTR is short in the Ensembl annotation. There is no peak in the DRS data at the end of either the RefSeq or Ensembl 3' UTR, but there are four peaks ~1.45, 1.9, 2.4 & 4.2 kb downstream of the existing Ensembl annotation (Figure 1, Track A, 1–4, respectively). These peaks all have canonical AATAAA poly(A) motifs (≤1 mismatch) located 15–22 bp upstream suggesting they are genuine poly(A) sites, however the DRS data alone do not reveal which, if any, of these sites should be associated with BMPR1A.

EST and RNA-seq data can provide a bridge between the Ensembl/RefSeq annotations and the DRS data. Despite their low depth, the G. gallus EST data show almost continuous coverage between the end of the 3' UTR annotated in Ensembl and the most 3' DRS peak. However, the EST data are not conclusive; there is a 400 bp gap in the EST coverage and the implied exon structure is inconsistent with the existing annotations. The addition of publicly available RNA-seq data ([24]) strengthens the confidence that the DRS peaks correspond to the 3'-end of BMPR1A. The RNA-seq data cover the proposed 3' UTR and finish 1 bp beyond the fourth DRS peak. The RNA-seq data also confirm the exon/intron structure of the existing gene annotations.

Although the RNA-seq data are non-uniformly distributed, there are only three places in the proposed 3' UTR where the read depth drops to zero. In all three examples, there is good supporting evidence from overlapping ESTs that these gaps are unlikely to represent the end of the gene. The combination of DRS, EST and RNA-seq suggests the BMPR1A gene in G. gallus should be re-annotated as shown in Table 1. The new annotation indicates four alternative poly(A) sites exist in the developing chicken embryo, but there is no evidence to support the two short novel protein coding models Ensembl also provide as annotations for this gene.

Complex, ambiguous, feature re-annotations: Chicken HOXA7. The re-annotation of BMPR1A was comparatively straightforward because the different datasets reinforce each other. A more complex and ambiguous re-annotation is illustrated in Figure 2 for the HOXA7 gene (ENSGALG0000011061, [33]). The Ensembl annotation has a single transcript that covers 1,792 bp and includes two exons (290 and 265 bp) and a short (36 bp) 3' UTR. In contrast, the RefSeq annotation covers 1,837 bp, includes three exons (276, 293 & 41 bp respectively) and has no defined 3' UTR.

The intron/exon structure of HOXA7 shown in Figure 2 appears to be simpler than BMPR1A. However, the DRS, EST and RNA-seq datasets suggest this gene may have a more complex structure than defined in Ensembl/RefSeq. Multiple peaks are evident in the observed DRS dataset (Figure 2, Track K, 1–6) that mark potential poly(A) sites associated with HOXA7. The first peak (1) lies within the intron separating the two primary exons of the gene. The second peak (2) is composed of three smaller peaks that all lie within 30 bp of the end of the existing Ensembl annotation. On the surface, these appear to support the existing 3' UTR annotation, but the presence of a large peak in the DRS data 1.5 kb downstream (6), if genuinely associated with HOXA7, suggests an alternative annotation that would not extend the 3' UTR, but would also be the dominant transcript in the DRS dataset for this gene. Peak 6 shows a canonical AATAAA poly(A) motif 19 bp upstream, consistent with a genuine poly(A) site. Peaks 2–5 show long runs of adenosine bases immediately downstream of each peak, suggesting that they might be the result of internal priming while peak 1 shows neither of these features and it remains unclear whether it is a true site of alternative polyadenylation.

In a similar fashion to the example shown in Figure 1 (Section 1.1), both the EST and RNA-seq data bridge the gap between DRS peak 6 and the existing reference annotations. Together, these data support the proposed 3' UTR re-annotation, despite the EST data including a 500 bp region where the coverage is low (≤2 ESTs) and from an inferred exon structure that is inconsistent with the existing annotation.

While the RNA-seq data support the proposed 3' UTR re-annotation, they do not match the short initial exon present in the RefSeq annotation and the EST data. The genomic sequence in the 31 bp intron between the first and second exons in the RefSeq annotation is marked as 'N's in the genomic sequence, making it difficult to draw robust conclusions on the structure of the gene in this region. Although this exon annotation is broadly supported by the EST dataset, these data extend beyond the RefSeq annotation suggesting a potential re-annotation of the 5' UTR.

This example shows considerable non-uniformity in the RNA-seq data that map to the suggested 3' UTR, with several significant (>50 bp) gaps in the RNA-seq coverage. The EST coverage and the lack of known polyadenylation motifs in the genomic sequence surrounding these gaps suggest that these are artefacts intrinsic to the Illumina RNA-seq protocol and do not represent the end of the 3' UTR associated with HOXA7.

Accordingly, a re-annotation of the HOXA7 gene in G. gallus (Table 2) based on the combination of DRS, EST and RNA-seq data is proposed. The annotation broadly supports the existing intron/exon structure of the RefSeq annotation, but extends the 3' UTR by 1.5 Kb and suggests an alternative polyadenylation site. The presence of the first intron is not strongly supported by the RNA-seq data and may well be spurious or an extension of the larger second exon, or specific to a particular tissue type or biological condition not sampled by the RNA-seq experiment.

Since completion of this study, the Galgal3 genome has been superseded by Galgal4 (released in Nov 2011) and its corresponding annotations (ensembl v71 and later, Apr 2013). Despite the undoubted improvements this new version has made to the genome as a whole, the gene models for both BMPR1A and HOXA7 have not changed significantly and our proposed...
reannotations for these genes remain pertinent for the latest Galgal4 annotations (ensembl v74).

Gene and 3' UTR re-annotation for Homo sapiens SLFN5. Although the human genome is actively curated, gene models can still be revised with new data. For example, SLFN5 in H. sapiens until recently had a significantly truncated 3' UTR. Prior to v69 (Oct 2012), the SLFN5 Ensembl annotation was composed of two alternative gene models; one covering 4,625 bp spanning 4 exons, and the other covering 2,540 bp spanning 3 exons. The RefSeq annotation contained a single gene model covering 4,654 bp spanning 4 exons. All these annotations included a short 5' UTR encompassing a long intron and a well-defined 1.8 kb 3' UTR. In the v69 Ensembl release, the annotations for SLFN5 changed considerably. The 3' UTR for the primary transcript was extended by ~6 kb and a third, shorter gene model was added. To date (Feb 2013), there has been no

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Figure 1. The genomic context around BMPR1A in G. gallus. Figures 1–8 are divided into three regions comprising information located on the forward strand (pink), reverse strand (grey) and un-stranded information (yellow). Each region is subdivided into tracks showing a selection of the different annotations/datasets described below. For clarity, tracks are omitted where the track contains no data in the region shown. Tracks A & K: Histograms for forward (A) and reverse (K) strands computed by summing the number of uniquely aligned DRS reads that end at a position and presented in units of read-counts/base. Tracks B & J: Filled rectangles show forward (B) and reverse (J) strand individual EST alignments for a selection of the total EST coverage. Individual EST alignments that span across an implied exon splice junction are illustrated by a split bar representing the sequenced EST joined by a thin line that spans the implied intron. Tracks C & I: Additional annotation information for forward (C) and reverse (I) strands. This track shows annotation information that doesn’t originate from a primary reference database for the species. Details of the specific annotations shown for each figure are given in the figure caption. Tracks D & H: Primary database annotations labelled with the database primary identifier for forward (D) and reverse (H) strands. Multiple gene models are shown where appropriate. Exons are shown as thick bars, UTRs as thinner bars and introns as thin lines. For A. thaliana this track shows the TAIR (v10) annotations. For the other examples in this paper, this track shows Ensembl (v69, red) and RefSeq (v191, green) annotations. Track E: Unstranded RNA-seq read depth histogram, computed by summing the number of uniquely aligned reads that cover at any given position and expressed in read counts/base. Track F: RNA-seq individual read alignments, for a selection of the total read depth, shown as filled rectangles. Individual read alignments that span across an implied exon splice junction are represented by a split bar representing the sequenced read joined by a thin line showing the implied intron. Track G: Unstranded sRNA-seq read depth histogram, computed by summing the number of uniquely aligned sRNA-seq reads that cover at any given position and expressed in units of read-counts/base. Figure 1 shows a ~57 kb region of G. gallus, chromosome 6, including BMPR1A (ENSGALG00000003119) and illustrates a straight-forward gene re-annotation, where the RNA-Seq and DRS data combined are sufficient to define the extent, structure, and alternative polyadenylation positions for a gene. Tracks C & I show confirmed complete coding sequence mRNA data for the region (GenBank v191 - orange) and the locations of the Affymetrix chicken GeneChip microarray probe-sets (black markers), and the cDNA against which the Affymetrix probe-sets were designed (light blue). See the Materials and Methods section for more details on the generation and processing of the G. gallus RNA-seq and DRS data-sets. The EST data (B & J) are from [47]. The DRS track for the reverse strand (Track H) contains no data in the region shown and has been removed for clarity.
Table 1. Comparison of annotations for BMPR1A.

| Primary annotation       | Chr | Begin (bp) | End (bp) | Strand | Coverage (bp) |
|--------------------------|-----|------------|----------|--------|---------------|
| RefSeq: BMPR1A           | 6   | 3,546,262  | 3,585,602 | +      | 39,340        |
| ensembl: ENSGALT0000003119 | 6   | 3,546,283  | 3,585,813 | +      | 39,530        |

Proposed re-annotation

| EST/RNA-seq: 5’ UTR   | 6   | 3,546,262  | 3,564,179 | +      | 17,917        |
| EST/RNA-seq: BMPR1A   | 6   | 3,564,180  | 3,585,585 | +      | 21,405        |
| DRS/EST/RNA-seq: 3’ UTR | 6   | 3,585,586  | 3,590,064 | +      | 4,478         |
| Summary                | 6   | 3,546,262  | 3,590,064 | +      | 43,800        |

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Figure 2. The genomic context around HOXA7 in G. gallus. The individual tracks and layout of this figure are as described in Figure 1. Figure 2 shows a ~6 kb region of G. gallus, chromosome 2 that encompasses HOXA7 gene. The RNA-seq (Tracks E & F), Helicos BioSciences’ DRS (Tracks A & K) and publically available EST (Tracks B & J) datasets for this region are ambiguous, but combined, the data clearly define the extent, and structure for this gene. Tracks C & I show the same additional annotation tracks as shown in Figure 1. See the Materials and Methods section for more details on the generation and processing of the G. gallus RNA-Seq and DRS data-sets. EST data were taken from [47].

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change in the RefSeq annotation for this gene. Figure 3 shows the genomic context around \textit{SLFN5} with the most recent annotations from Ensembl and RefSeq. Both the DRS and RNA-seq data show evidence for transcription continuing up to \( \approx 6 \) kb further downstream than the current RefSeq annotation, and in agreement with the current Ensembl annotation. However, the DRS data reveals two alternative polyadenylation sites \( \approx 5 \) kb and \( \approx 8.5 \) kb (Figure 3, Track A, 1--2, respectively) from the first stop codon in \textit{SLFN5}, both of which have the canonical AATAAA cleavage and polyadenylation signal upstream (19 & 24 bases, respectively) of the DRS peak. One of these sites is coincident with the Ensembl gene model, but the second site suggests a fourth alternative gene model. The combination of the DRS and RNA-seq data provides a more complete picture of the splice variants and transcriptional boundaries of the \textit{SLFN5} gene.

| Table 2. Comparison of annotations for \textit{HOXA7}. |

| Primary annotation     | Chr | Start (bp) | Stop (bp) | Strand | Coverage (bp) |
|------------------------|-----|------------|-----------|--------|---------------|
| RefSeq: \textit{HOXA7} | 2   | 32,570,322 | 32,572,159| -      | 1,837         |
| ensembl: ENSGALT00000018013 | 2   | 32,570,285 | 32,571,987| -      | 1,702         |

| Proposed re-annotation |
|------------------------|
| EST/RNA-seq: 5' UTR    | 2   | 32,572,160 | 32,572,292| -      | 132           |
| EST/RNA-seq: \textit{HOXA7} | 2   | 32,570,322 | 32,572,159| -      | 1,837         |
| DRS/EST/RNA-seq: 3' UTR | 2   | 32,568,768 | 32,570,321| -      | 1,553         |
| Summary                | 2   | 32,572,160 | 32,570,321| -      | 3,522         |

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Figure 3. The genomic context around \textit{SLFN5} in \textit{H. sapiens}. This figure shows a \( \approx 6 \) kb region of \textit{H. sapiens}, chromosome 17, that encompasses the newly re-annotated \textit{SLFN5} gene. Two peaks in the DRS data for this region (Track A) reveal that even our most up-to-date annotations in heavily curated genomes are often incomplete. The difference between the annotations provided by RefSeq and Ensembl (Track D) also highlights that existing primary database annotations often disagree significantly, making downstream analysis results dependent of the reference database used for individual studies. For full details of the individual tracks and layout of this figure, see the legend to Figure 1. See the Materials and Methods section for more details on the generation and processing of the \textit{H. sapiens} RNA-seq and DRS data-sets.

doi:10.1371/journal.pone.0094270.g003
The RNA-seq data suggests the SLFN5 gene in \textit{H. sapiens} should be re-annotated as described in Table 3.

\textbf{Extension of 3' UTR for \textit{A. thaliana}: AT4G02715.} The genome of \textit{A. thaliana} has been extensively studied since it was sequenced and released in 2000 ([34]). However, examination of the first DRS data for \textit{A. thaliana} [19] enabled the 3'-ends of \textasciitilde65% of its genes to be re-annotated automatically by considering reads within 300 bp of the TAIR10 annotated 3'-end. Sherstnev et al. [19] only considered DRS data and this approach missed further re-annotation possibilities. For example, Figure 4 summarises the region around \textit{AT4G02715}. The TAIR10 annotation for this gene consists of a 0.6 kb 5'-UTR containing a single intron followed by a single 0.6 kb exon. No significant DRS peaks are found within the 300 bp window downstream of the 3' end of the current annotation and so the algorithm described in [19] did not re-annotate the 3' end of this gene. A cluster of DRS signals is observed \textasciitilde0.6 kb downstream (Figure 4, Track K, 2) followed by a set of peaks \textasciitilde0.65 kb further downstream (Figure 4, Track K, 3) and another cluster of peaks \textasciitilde0.25 kb still further downstream (Figure 4, Track K). The RNA-seq data covers the full extent of the downstream region up to DRS peak 3. Like many poly(A) sites in \textit{Arabidopsis}, peak 3 is composed of at least four peaks of varying strength, several of which are broader than the \pm2 bp positional accuracy of the DRS data [19]. The RNA-seq data also identify an intron \textasciitilde1 kb upstream of the end of the current annotation. The protein coded by AT4G02715 has yet to be characterized and the current annotation represents the longest ORF in this genomic region, suggesting that the proposed extension reflects the 3' UTR of this gene. The RNA-seq data show weak expression extending out to within a few bases of peak 4, but the unmatched nature of the DRS and RNA-seq samples makes it difficult to draw strong conclusions about the nature of this region. It is possible this region is an alternative transcript for AT4G02715 that is not expressed in the archival RNA-seq dataset.

Table 4 shows the proposed re-annotation of AT4G02715 in \textit{A. thaliana} based on the RNA-seq and DRS data. In the new annotation, the DRS data describes the primary gene transcript and tentatively suggests the presence of alternative transcripts.

\textit{A. thaliana}: AT1G68945 – annotation and data inconsistent. Figure 5 shows AT1G68945 which has been confirmed as protein coding from cDNA and EST data, although the protein product has yet to be characterized. It has only one annotated gene model, comprising a long 5' UTR, a single coding exon, and a short 3' UTR. No significant DRS peaks are found associated with this gene model or within the 300 bp window downstream of the 3' end of the current annotation and so the algorithm described in [19] does not re-annotate this gene and leads to the conclusion that it is not expressed. Curiously however, a strong signal is seen in the DRS data on the opposite strand, at the start of the 5' UTR annotation. This peak is broad, covering \textasciitilde20 bp, suggesting multiple possible poly(A) sites. Reads from the un-stranded RNA-seq data align precisely to the gene position confirming its location but not which strand it is on. One possible interpretation of this region is that there is a gene on the reverse strand that is not annotated in TAIR10 (as suggested in Table 5) this is also supported by single-stranded RNA-Seq data from the Ecker Lab [35]. However, the reverse strand in this region of the current genome build contains multiple stop codons suggesting it is unlikely to represent a single protein coding gene.

\textbf{Section 2: Disentangling gene expression in complex genomic regions}

\textit{Homo sapiens}: Mettl12. Figure 6 illustrates the genomic region around the gene Mettl12 which is located on the forward strand of chromosome 11. This region shows the challenges of annotation and expression quantification in complex regions and how combining different datasets, in particular strand-specific data that defines 3'-ends, can help alleviate some of these difficulties. Ensembl v69 provides several different gene annotation models for Mettl12, while RefSeq reports a single gene model that is significantly different to the Ensembl annotations. All these models agree on a 5' UTR that includes an intron, within which resides a copy of the snoRNA, snorna57 (this is one of four copies of this snoRNA that occur in the human genome). The Mettl12 locus is additionally complicated by the presence of a large protein-coding ORF, C11orf48, on the antisense strand that overlaps Mettl12 completely. Ensembl provides a total of thirteen different gene models for C11orf48, while RefSeq lists a single gene model. In addition, the annotated 5' UTRs of several C11orf48 gene models overlap with the 5' UTR of the forward strand ORF C11orf83,
which itself has two separate gene models. The details of all these annotations are provided in Table 6.

As one might anticipate for such a complex region, the unstranded Illumina RNA-seq data for this region are ambiguous, so quantifying gene expression from these data is problematic. The terminal four exons of \textit{C11orf48} are strongly-expressed (read depth, 150–300) suggesting that the gene model \textit{ENST00000524958} is the predominant expressed form of \textit{C11orf48} in these data. This is reinforced by reads that map across the intron/exon boundaries for this gene model. Importantly, there are no reads mapping across any splice junctions immediately prior to the start of this annotation, clearly delineating this model from the others for \textit{C11orf48}. Similarly, two exons of \textit{C11orf83} are also strongly-expressed and show a consistent splicing pattern, but the expression appears to be truncated at a position that is inconsistent with all the current 3' UTR annotations for \textit{C11orf83}, suggesting a possible new gene model for this gene. The picture in the intervening region, which covers \textit{Mettl12}, \textit{snorna57} and another gene model for \textit{C11orf48}, is far less clear. The low-level expression in this region shows little in the way of distinct exon/intron boundaries that would help to identify the origin for this expression, but marginal evidence for some other transcripts of \textit{C11orf48} and for expression from \textit{Mettl12} can be identified from individual reads that map across appropriate exon/intron boundaries.

In contrast, the DRS data are more straightforward to interpret and quantify, since they reliably identify the sequenced strand. Hence, they can be used to help inform the gene annotations and quantify the gene expression in human skin within this genomic region. The DRS data have four distinct sites of expression; three on the forward strand (Figure 6, Track A, 1–3) and one on the reverse strand (Figure 6, Track K). On the forward strand, peaks 1 & 2 coincide with the \textit{Mettl12} annotations. Peak 1 is located in the 5' UTR of the annotations but downstream of \textit{snorna57} suggesting that this peak represents expression of the snoRNA precursor rather than the gene. Peak 2 is located in the annotated 3' UTR of \textit{Mettl12}, however it is only 13 bp downstream of the stop codon. The sequence in this region does not show any strong candidates for internal priming and the upstream sequence contains a slight variation on the canonical poly(A) motif (ATTAAA) 17 bp upstream. Although this signal hints at a new gene model for \textit{Mettl12}, with a short 3' UTR, the low-level of the expression

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\textbf{Figure 4. The genomic context around AT4G02715 in A. thaliana.} A ∼3 kb region of \textit{A. thaliana} on chromosome 4 is shown, which encompasses \textit{AT4G02715}. In this case the extensive 3' UTR extension suggested by the DRS data (Track K) shows how this re-annotation was missed even by the automated re-annotation algorithm applied in [19]. For full details of the individual tracks and layout of this figure, see Legend to Figure 1. See the Materials and Methods section for more details on the \textit{A. thaliana} RNA-seq, EST and DRS data-sets, and their processing. doi:10.1371/journal.pone.0094270.g004
Table 4. Comparison of annotations for AT4G02715 gene locus.

| Primary annotation | Chr | Start (bp) | End (bp) | Strand | Coverage (bp) |
|--------------------|-----|------------|----------|--------|---------------|
| TAIR10: AT4G02715  | 4   | 1,203,279  | 1,202,169| -      | 1,110         |
| Proposed re-annotation 1 | RNA-seq/EST: 5' UTR | 4 | 1,203,279 | 1,202,998 | - | 281 |
| | RNA-seq/EST: AT4G02715 | 4 | 1,202,998 | 1,202,169 | - | 829 |
| | RNA-seq/DRS/EST: 3' UTR | 4 | 1,202,169 | 1,200,886–1,200,975 | - | 1,194–1,279 |
| Summary | 4 | 1,203,279 | 1,200,886–1,200,975 | - | 2,304–2,389 |
| Proposed re-annotation 2 | RNA-seq/EST: 5' UTR | 4 | 1,203,279 | 1,202,998 | - | 281 |
| | RNA-seq/EST: AT4G02715 | 4 | 1,202,998 | 1,202,169 | - | 829 |
| | RNA-seq/DRS/EST: 3' UTR | 4 | 1,202,169 | 1,200,688 | - | 1,481 |
| Summary | 4 | 1,203,279 | 1,200,688 | - | 2,591 |
| Proposed re-annotation 3 | RNA-seq/EST: 5' UTR | 4 | 1,203,279 | 1,202,998 | - | 281 |
| | RNA-seq/EST: AT4G02715 | 4 | 1,202,998 | 1,202,169 | - | 829 |
| | RNA-seq/DRS/EST: 3' UTR | 4 | 1,202,169 | 1,200,666 | - | 1,503 |
| Summary | 4 | 1,203,279 | 1,200,666 | - | 2,613 |

doi:10.1371/journal.pone.0094270.t004

Figure 5. The genomic context around AT1G68945 in A. thaliana. This figure shows a ~600 bp region of A. thaliana, chromosome 1, around the existing annotation of the gene AT1G68945. In this case, the DRS data for this region (Track K) reveal that the existing annotation is on the incorrect strand. This kind of situation is difficult for automated re-annotation pipelines to deal with, particularly if they focus on using natively unstranded data, such as Illumina RNA-Seq, to inform the annotation. This highlights necessity of natively stranded data, such as DRS data, for correctly defining feature annotations. For full details of the individual tracks and layout of this figure, see Figure 1 (caption). See the Materials and Methods section for more details on the A. thaliana RNA-Seq, EST and DRS datasets, and their processing.

doi:10.1371/journal.pone.0094270.g005
makes this inconclusive. Further downstream on the forward strand, \textit{C11orf83} is strongly expressed in the DRS data (peak 3), again with an apparently shorter 3' UTR than annotated. The details of all these novel transcript annotations are provided in Table 6. The data are not as clear for the reverse strand. Assuming the current annotations are correct, the exquisite positional precision of the DRS data and the lack of any DRS peaks at other locations on the reverse strand, suggest four strong gene-model candidates. Of these, model \textit{ENST00000524958} is consistent with the DRS data and the RNA-seq data, supporting the conclusion that this is the predominantly expressed form of \textit{C11orf48}, at least in these samples. The other potential models may be correct, just not expressed in these samples.

**Homo sapiens: RPL31.** Figure 7 illustrates another example of a complex genomic region with ambiguous expression for convergent genes on opposite strands. On the forward strand, \textit{RPL31} has eleven gene models annotated in Ensembl, and three annotated in RefSeq. Across these models, the 5' UTR is annotated with seven different start positions and the 3' UTR is annotated with twelve alternative end positions. On the reverse strand, \textit{TBC1D8} is similarly complex, with ten Ensembl gene models and one RefSeq, four of which overlap with the five longest forms of \textit{RPL31}.

Again, as one might expect for such a complex region, the RNA-seq data are ambiguous. The RNA-seq data include five strong peaks that echo the exon and UTR structure of five of the \textit{RPL31} gene models, however the considerable low-level expression covering much of the region makes it hard to draw firm conclusions from RNA-seq data alone. This ambiguity is dramatically reduced with the addition of the DRS data. In the DRS data, a strong signal is observed coincident with the downstream edge of the fifth RNA-seq peak (Figure 7, Track A, Figure 7).

### Table 5. Comparison of annotations for \textit{AT1G68945} gene locus.

| Primary annotation | Chr | Start (bp) | End (bp) | Strand | Coverage (bp) |
|--------------------|-----|------------|----------|--------|---------------|
| TAIR10: \textit{AT1G68945} | 1 | 25,926,962 | 25,927,330 | + | 368 |
| **Proposed re-annotation** | | | | | |
| RNA-seq/EST: 5' UTR | 1 | 25,927,329 | 25,927,314 | - | 15 |
| RNA-seq/EST: \textit{AT1G68945} | 1 | 25,927,313 | 25,927,167 | - | 146 |
| RNA-seq/DRS/EST: 3' UTR | 1 | 25,927,166 | 25,926,947–25,926,967 | - | 199–219 |
| **Summary** | 1 | 25,927,329 | 25,926,947–25,926,967 | - | 360–380 |

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### Table 6. Comparison of annotations for \textit{Mettl12} gene locus.

| Primary annotation | Chr | Start (bp) | End (bp) | Strand | Coverage (bp) |
|--------------------|-----|------------|----------|--------|---------------|
| RefSeq: \textit{Mettl12} | 11 | 62,432,779 | 62,434,923 | + | 2,145 |
| \textit{ensembl}: \textit{ENST00000532971} | 11 | 62,432,781 | 62,435,580 | + | 2,800 |
| \textit{ensembl}: \textit{ENST00000398922} | 11 | 62,432,781 | 62,434,869 | + | 2,089 |
| \textit{ensembl}: \textit{ENST00000529868} | 11 | 62,432,785 | 62,435,968 | + | 3,184 |
| **Proposed re-annotation** | | | | | |
| RNA-seq: 5' UTR | 11 | 62,432,794 | 62,433,350 | + | 557 |
| RNA-seq: \textit{Mettl12} | 11 | 62,433,351 | 62,434,522 | + | 1,172 |
| RNA-seq/DRS: 3' UTR 1 | 11 | 62,433,867 | 62,434,535 | + | 4,992 |
| **Primary annotation** | | | | | |
| RefSeq: \textit{snoRNA57} | 11 | 62,432,893 | 62,433,041 | + | 148 |
| \textit{Ensemble}: \textit{ENST00000206597} | 11 | 62,432,893 | 62,433,041 | + | 149 |
| **Additional annotation** | | | | | |
| \textit{snoRNA57 precursor} | 11 | 62,432,794 | 62,433,179 | + | 385 |
| **Primary annotation** | | | | | |
| RefSeq: \textit{C11orf83} | 11 | 62,439,125 | 62,441,161 | + | 2,036 |
| \textit{ensembl}: \textit{ENST00000531323} | 11 | 62,437,745 | 62,441,049 | + | 3,304 |
| \textit{ensembl}: \textit{ENST00000377953} | 11 | 62,439,126 | 62,441,159 | + | 2,033 |
| **Proposed re-annotation** | | | | | |
| RNA-seq: 5' UTR | 11 | 62,439,125 | 62,439,216 | + | 91 |
| RNA-seq: \textit{C11orf83} | 11 | 62,439,217 | 62,439,584 | + | 367 |
| RNA-seq/DRS: 3' UTR 1 | 11 | 62,439,585 | 62,439,844 | + | 259 |
| **Summary** | 11 | 62,439,125 | 62,439,844 | + | 719 |

doi:10.1371/journal.pone.0094270.t006
peak 1). This broad peak covers ~20 bp and encompasses the 3' UTR ends of seven of the annotated models. The sequence immediately upstream of the peak is strongly AT rich, suggesting that the location of the poly(A) site in RPL31 may not be very precisely controlled. Instead, a range of possible poly(A) positions occur with different likelihoods within this window.

Interestingly, two small peaks also occur in the DRS data further downstream (Figure 7, Track A, peaks 2 & 3), close to three of the longer RPL31 gene models. The first of these extends the nearest gene model by 56 bp, the second lies within 5 bp of the end of the longest annotated model. Both of these peaks have the AATAAG variant of the canonical polyadenylation signal, ~19 bases upstream of the peak. This weak but distinct signal clearly demonstrates that the shorter RPL31 gene models are not the only form of transcripts made from this gene in these data. On the reverse strand, the DRS data shows a strongly expressed peak...
(Peak 5) that is coincident with the end of the 3' UTR annotated in the RefSeq TBC1D8 gene model. However, a second peak ~1.2 kb further downstream (Peak 4), identifies a new putative polyadenylation site for this gene. Both of these peaks show the polyadenylation motif AATAAG ~20 bp upstream. Accordingly, a new gene model is proposed for RPL31 that results in a transcript that overlaps with all the RPL31 gene models. The details are highlighted in Table 7.

Section 3: A clearer picture of small RNA expression

It is currently not possible to quantify the expression of long and short RNAs in a single RNA-Seq experiment. In order to identify expression of mature miRNAs, in particular, a protocol is used that specifically selects very short (<30 bp) RNA species and so excludes the ~200 bp fragments commonly selected by RNA-seq protocols. Mature intergenic miRNAs are ~21 bp single stranded RNA molecules processed out of pre-miRNA hairpin loops found in pri-miRNA transcripts and are transcribed by RNA polymerase II ([36]). The pri-miRNAs have been shown to be polyadenylated.

Figure 7. The genomic context around RPL31 in H. sapiens. This ~25 kb region of H. sapiens, chromosome 2, again highlights the difficulties in interpreting unstranded data in complex genomes. This region encompasses the gene RPL31 on the forward strand and TBC1D8 on the reverse strand. Many of the existing annotations for these two genes overlap (Tracks D & H) making unstranded data difficult to interpret with certainty. The natively stranded DRS data (Tracks A & K) clearly delineate the ends of the transcripts observed from both these genes, including a new annotation for TBC1D8. For full details of the individual tracks and layout of this figure, see Figure 1 (caption). See the Materials and Methods section for more details on the H. sapiens RNA-seq and DRS datasets, and their processing. doi:10.1371/journal.pone.0094270.g007
via a variety of methods including PCR primers ([36–38]), sequence analysis ([39]) and sequencing ([40]). The miRNA*, which is not loaded in the RISC complex is not normally retained, but can often be observed in high-throughput sequencing.

miR-200c and miR-141 illustrate the advantages of combining DRS and RNA-seq data with small RNA-seq (sRNA-seq) data for a better characterisation of intergenic pri-miRNAs. Figure 8 shows the genomic region around miR-200c and miR141. This region is flanked by genes that are expressed in the DRS and RNA-seq data; PTPTN6 on the forward strand (Figure 8, Track D) and PHB2 on the reverse strand (Figure 8, Track H). Aligning directly with the miRNA annotations are two pairs of peaks in the sRNA-seq data (Figure 8, Track G, 1 & 2) that correspond to the mature miRNAs miR-200c-5p/3p and miR-141-5p/3p sequences. In each case, the 3p sequences are the dominant expressed form, as shown by the relative heights of the sRNA-seq peaks within each pair.

The structure and extent of the pri-miRNA is clearly delineated by the RNA-seq data (Figure 8, Track E) in the regions flanking the two mature miRNA loci. No reads are detected within the intronic region that covers the pre- or mature miRNA regions suggesting that the pre-miRNAs processing and cleavage occurs rapidly, leaving the 5’ and (polyadenylated) 3’ end fragments to be slowly degraded. The DRS data support this picture showing a cluster of expression, 200 bp downstream of miR-141-3p on the forward strand (Figure 8, Track A, 3) that has the tandem polyadenylation site motif AATAAATAAA 26 bp upstream.
Section 4: Novel gene discovery

In addition to improving existing annotations, the combination of DRS, RNA-seq and other datasets also identifies and characterises genomic regions containing new feature candidates. The discovery of potential new snoRNAs in the downstream region of the gene AT4G10810 in A. thaliana, shown in Figure 9, is an example. The RNA-seq data downstream of AT4G10810 shows significant low-level expression over a ~600 bp region, with no strong evidence for intron/exon structure (Figure 9, Track E, 1). The DRS data in this region are complex, showing a considerable number of small peaks that suggest multiple possible alternative polyadenylation sites (Figure 9, Track K, 2). Combined, these imply a cluster of short, currently un-annotated, features. This picture is reinforced by the large peak in expression seen in the sRNA-seq data in this region (Figure 9, Track G, 3). This peak does not show the two-peak structure characteristic of mature miRNA sequences (see Section 4), leaving us to speculate on the nature of this short feature. The SnoSeeker (v1.1, [41]) snoRNA prediction algorithm predicts a snoRNA coincident with this
position suggesting that this is a previously undiscovered snoRNA gene. The details for the novel gene structure are in Table 8.

**Discussion**

Detailed, complete, genomic feature annotations are a cornerstone of modern biology. Their importance, particularly for experiments that rely on high-throughput transcriptomics, cannot be overstated. However, defining these annotations is not a trivial task and is made more difficult by the fact that there may be multiple `correct` annotations for a gene. While the importance of accurate annotations is widely recognised, the impact that alternative individual annotation, or an alternative set of annotations, has on the subsequent downstream analysis (e.g.,

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**Figure 9. The genomic context around AT4G10810 in A. thaliana.** This figure shows a ~2 kb region of A. thaliana, chromosome 4, including AT4G10810 that demonstrates the capability of combined DRS, RNA-seq and sRNA-seq to identify novel genes. This also highlights some of the limitations of automated re-annotation algorithms that are based on arbitrarily chosen parameter values. In this case, [19] (2012), provide a re-annotation of the 3' UTR of AT4G10810 by focussing on the DRS data within a region 300 bp downstream of the end of the primary database annotations (Track K). For most A. thaliana genes, this proves to be an effective strategy, but occasionally it results in incorrect re-annotations. Here, the region downstream of AT4G10810 encompasses multiple relatively weak DRS peaks (Track K, 2) and Sherstnev et al mistakenly re-annotate the gene to include many of these peaks (Track I). In fact, the RNA-seq data (Tracks E & F, 1) clearly identify the spatial separation between AT4G10810 and the significant low-level downstream expression, suggesting a novel gene, or cluster of genes. Interestingly, a strong peak in the sRNA-seq data in this region (Track G, 3), coupled with a coincident prediction from SnoSeeker (Track I), strongly suggests the presence of a novel snoRNA in this region. See the **Materials and Methods section** for more details on the generation and processing of the A. thaliana RNA-seq, sRNA-seq, EST and DRS datasets.

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differential gene expression) and biological understanding is less well appreciated. Two distinct classes of problem occur commonly for genome annotations; an incomplete set of feature annotations and/or an unreliable individual feature annotation.

The known set of human genes is an example of an incomplete set of feature annotations, i.e. a set of individual annotations (each of which may also be complete), that is missing discrete members of the set. Over the past decade considerable effort has been expended in manually curating the annotations for the human genome. As a consequence, the annotations for known genes is precise given the available data but the set as a whole is still likely to be missing as yet undiscovered genes and alternatively processed mRNA isoforms ([29]). For human and other heavily curated genomes, even though the full set of information is not known, the information that exists for the individual annotations is often reliable. Providing the set is not too incomplete, this will have relatively little impact on downstream analyses that rely on these annotations. One important exception is where features that are not annotated overlap completely with known features. For example, the observed fold change for such a region could be completely misleading and would not reflect the underlying biology if expression of the overlapping genes is very different.

Unreliable individual annotations present a different challenge. Here, members of a set of feature annotations (that may be partially complete) are based on a limited or significantly imprecise set of information. The impact this has on any downstream data analyses depends on the properties of the data being used and the specific analyses. For example, differential gene expression between two experimental conditions based on RNA-seq data is not dramatically sensitive to having marginally inaccurate annotation of gene structure unless the gene structure changes between conditions. Since the conditions being compared both use the same annotations, and given that the annotations are covered by a significant majority of the reads, the calculated fold change will be similar to the actual fold change that would be calculated using a more accurate set of annotations. Techniques that focus on one region of the gene such as DRS are far more sensitive to inaccurate or incomplete annotation information. If the locus that has been sequenced is not included within the annotated GARs of the gene then no (or very little) expression will be attributed to this gene in either condition, regardless of the true change in expression in the data.

For most published genomes, the available annotation is the result of an automated prediction-based annotation pipelines (see, for example, [42], [43]). Automated gene prediction is a difficult challenge (see [44]) and these first-pass annotations often contain considerable inaccuracies. Re-annotation using automatic methods typically involves discarding the current set of annotations and building the annotations again from scratch as the genome sequence is improved. In some cases, re-annotation has been attempted by supplementing the current annotations guided by high-throughput transcriptomics sequencing data ([19]). Automated, but data-driven, re-annotations can provide a considerable increase in the quality of feature annotations however they still have several drawbacks. Typically automatic methods depend on several arbitrarily set parameters such as the size of the window probed for new feature endpoints and the minimum number of reads required to extend an annotation (this is also true of automated annotation pipelines). As a result, many individual feature annotations will remain inaccurate and/or the annotation set remain incomplete. The A. thaliana re-annotation provided by [19] considerably extends and improves on an already comprehensive and detailed genome annotation in a well-studied model species (TAIR version 10 - [45]). However, the automated annotation method is unable successfully to re-annotate genes requiring a 3′ extension longer than the 300 bp downstream window, nor can it distinguish between a genuine new 3′ end annotation or the 3′ end of a new short gene located immediately downstream of an existing annotation (see, for example, Section 5 and Figure 9). Even after re-annotation dozens of intergenic DRS peaks (many comprised of >50 raw reads) remain un-accounted for, indicating the need for a more careful data-driven re-annotation.

The majority of high-throughput transcriptomics sequencing datasets are not generated with the primary intention of re-annotating genomic features, yet these datasets provide a wealth of information that can do exactly that. Individual sequencing technologies often show characteristics that make it difficult to base strong conclusions about feature re-annotation solely on the data they generate (Table 9). The experience gained in the present study suggest that genome annotation efforts that focus on using a single data type (for example, [46]) are likely to have difficulty producing a high-quality, high-completeness set of feature annotations for eukaryotic genomes. Combining the strengths of RNA-seq data, short RNA-seq, archival EST/mRNA data and strand-specific sequencing that defines the 3′-end is particularly effective at overcoming the weaknesses inherent to data generated from any one of these technologies individually (Table 5). These data can be used to identify and characterise gene intron/exon structure, and characterise GARs associated with these genes. The

| Table 8. Transcript annotations for AT4G10810 gene locus. |
|----------------------------------------------------------|
| **Primary annotation**         | **Chr** | **Start (bp)** | **End (bp)** | **Strand** | **Coverage (bp)** |
|-------------------------------|---------|----------------|--------------|------------|------------------|
| TAIR10: AT4G10810             | 4       | 6,646,335      | 6,645,715    | -          | 620              |
| [19]                          | 4       | 6,646,335      | 6,645,421    | -          | 914              |
| **Proposed re-annotation 1**  |         |                |              |            |                  |
| RNA-seq/EST: 5′ UTR           | 4       | 6,646,335      | 6,646,229    | -          | 106              |
| RNA-seq/EST: AT4G6894S        | 4       | 6,646,230      | 6,645,984    | -          | 246              |
| RNA-seq/DRS/EST: 3′ UTR       | 4       | 6,645,985      | 6,645,715–6,645,864 | - | 121–270          |
| Summary                       | 4       | 6,646,335      | 6,645,715–6,645,864 | - | 471–620 |
| **Proposed re-annotation 2**  |         |                |              |            |                  |
| Novel snoRNA                  | 4       | 6,645,422      | 6,645,529    | -          | 107              |
| snoSeeker Predicted snoRNA    | 4       | 6,645,420      | 6,645,538    | -          | 118              |

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examples of gene re-annotation described here are the result of manual interpretation and integration of these different data. The steps in this manual process are not hard-and-fast rules, but rather flexible, somewhat fuzzy, interpretive decisions. A simplified flow diagram capturing the core steps of this process is shown in Figure 10 and highlights those aspects of this decision-making process that require qualitative judgement. While this flow diagram has proved difficult to automate in software, the framework may form the basis for the development of a logic-based system for re-annotation, or a rule-based “expert system” to help a skilled genome annotator. The DRS data is particularly important in this process, both by providing precise information about the termination point of 3’ UTRS and by unambiguously identifying the strand for the gene expression data. Accurately constraining 3’ UTRS associated with genes is particularly important for alternative polyadenylation studies, microRNA and other regulatory element binding site identification. It is also important for downstream differential gene expression analysis and functional pathway analysis, because a significant fraction of RNA-seq reads, and all DRS reads, associated with a gene lie within their associated 3’ UTR.

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Figure 10. Reannotation flow diagram. This flow diagram represents a distillation of the key aspects of the manual re-annotation process used for the examples presented here. Starting with loading the data into a genome browser (grey box, rounded corners), the process is a complex decision tree with several key stages that require judgement, experience and familiarity with the data in addition to quantitative information (blue boxes). This dependence on qualitative judgements makes this process extremely difficult to capture computationally (in addition to the fact that the process would hardly change if used with different datasets, species, etc), underscoring the importance of manual annotation. Paths through the decision tree end in one of eight possible end-points; three ‘positive’ re-annotation endpoints (green boxes, rounded corners), three ‘tentative’ re-annotation endpoints (orange boxes, rounded corners) and two ‘negative’ no re-annotation endpoints. Here we briefly look at the path through the decision tree for two of the simpler examples presented earlier in this work: Example Path 1: BMPR1A (Section 1, Figure 1) Starting with loading the data for BMPR1A (1), the EST, cDNA and RNA-seq support the existing annotation intron/exon structure (2), DRS peaks exist downstream (3), RNA-seq and EST data extend beyond the existing annotation (4), the EST and RNA-seq data terminate almost exactly coincident with the strongest downstream DRS peak (5), the EST and RNA-seq data are continuous coverage over the proposed extension (6), there are no clear sequence features (stop codon or internal priming signatures that strongly suggest the re-annotation would be incorrect (7), we propose a clear re-annotation of the gene (8). Example path 2: AT1G68945 (Section 1, Figure 5) Starting with loading the data for AT1G68945 (1), the EST, cDNA and RNA-seq support the existing annotation intron/exon structure (2), DRS peaks do not exist downstream (3), the EST and RNA-seq data are stranded, however they strand association is unreliable (9), there are sequence features (in this case, numerous stop codons in multiple frames (11), the data, sequence features and existing annotation are inconsistent. We cannot re-annotate the gene without more evidence.(12). In this case, further evidence in the form of strand-specific RNA-seq data from the Ecker Lab [35] would, if included, allow us to follow the path 1,2,3,9,10,13,14,15 resulting in a tentative re-annotation to the opposite strand, despite the apparent presence of stop codons.
automatic annotation pipelines will improve with the inclusion of strand-specific RNA-seq data and data that delineates the 5' and 3' ends precisely. Indeed, major projects such as Ensembl are now incorporating these data into their annotation pipelines (S. Searle per. Comm.). However, the examples presented in this paper suggest that for complete and precise annotation there is currently no substitute for annotation curated by experienced and knowledgeable scientists from a combination of DRS, RNA-seq, sRNA-seq, EST and other informative data.

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Author Contributions

Conceived and designed the experiments: NS CC KGS SJB GGS GJB. Performed the experiments: JS CD SJB. Analyzed the data: NS CC AS CD KGS SJB GGS GJB. Contributed reagents/materials/analysis tools: NS CC AS. Wrote the paper: NS CC AS CD KGS SJB GGS GJB. Contributed to discussions of the human skin data study and analysis: WHIM.

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