

INNOVATION

RNA-Seq: a revolutionary tool for transcriptomics

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Abstract | RNA-Seq is a recently developed approach to transcriptome profiling that uses deep-sequencing technologies. Studies using this method have already altered our view of the extent and complexity of eukaryotic transcriptomes. RNA-Seq also provides a far more precise measurement of levels of transcripts and their isoforms than other methods. This article describes the RNA-Seq approach, the challenges associated with its application, and the advances made so far in characterizing several eukaryote transcriptomes.

The transcriptome is the complete set of transcripts in a cell, and their quantity, for a specific developmental stage or physiological condition. Understanding the transcriptome is essential for interpreting the functional elements of the genome and revealing the molecular constituents of cells and tissues, and also for understanding development and disease. The key aims of transcriptomics are: to catalogue all species of transcript, including mRNAs, non-coding RNAs and small RNAs; to determine the transcriptional structure of genes, in terms of their start sites, 5' and 3' ends, splicing patterns and other post-transcriptional modifications; and to quantify the changing expression levels of each transcript during development and under different conditions.

Various technologies have been developed to deduce and quantify the transcriptome, including hybridization- or sequence-based approaches. Hybridization-based approaches typically involve incubating fluorescently labelled cDNA with custom-made microarrays or commercial high-density oligo microarrays. Specialized microarrays have also been designed; for example, arrays with probes spanning exon junctions can be used to detect and quantify distinct spliced isoforms¹. Genomic tiling microarrays that represent the genome at high density have been constructed and allow

the mapping of transcribed regions to a very high resolution, from several base pairs to ~100 bp^{2–5}. Hybridization-based approaches are high throughput and relatively inexpensive, except for high-resolution tiling arrays that interrogate large genomes. However, these methods have several limitations, which include: reliance upon existing knowledge about genome sequence; high background levels owing to cross-hybridization^{6,7}; and a limited dynamic range of detection owing to both background and saturation of signals. Moreover, comparing expression levels across different experiments is often difficult and can require complicated normalization methods.

“RNA-Seq [...] is expected to revolutionize the manner in which eukaryotic transcriptomes are analysed.”

In contrast to microarray methods, sequence-based approaches directly determine the cDNA sequence. Initially, Sanger sequencing of cDNA or EST libraries^{8,9} was used, but this approach is relatively low throughput, expensive and generally not quantitative. Tag-based methods were developed to overcome these limitations,

including serial analysis of gene expression (SAGE)^{10,11}, cap analysis of gene expression (CAGE)^{12–14} and massively parallel signature sequencing (MPSS)^{15–17}. These tag-based sequencing approaches are high throughput and can provide precise, ‘digital’ gene expression levels. However, most are based on expensive Sanger sequencing technology, and a significant portion of the short tags cannot be uniquely mapped to the reference genome. Moreover, only a portion of the transcript is analysed and isoforms are generally indistinguishable from each other. These disadvantages limit the use of traditional sequencing technology in annotating the structure of transcriptomes.

Recently, the development of novel high-throughput DNA sequencing methods has provided a new method for both mapping and quantifying transcriptomes. This method, termed RNA-Seq (RNA sequencing), has clear advantages over existing approaches and is expected to revolutionize the manner in which eukaryotic transcriptomes are analysed. It has already been applied to *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, *Arabidopsis thaliana*, mouse and human cells^{18–24}. Here, we explain how RNA-Seq works, discuss its challenges and provide an overview of studies that have used this approach, which have already begun to change our view of eukaryotic transcriptomes.

RNA-Seq technology and benefits

RNA-Seq uses recently developed deep-sequencing technologies. In general, a population of RNA (total or fractionated, such as poly(A)+) is converted to a library of cDNA fragments with adaptors attached to one or both ends (FIG. 1). Each molecule, with or without amplification, is then sequenced in a high-throughput manner to obtain short sequences from one end (single-end sequencing) or both ends (pair-end sequencing). The reads are typically 30–400 bp, depending on the DNA-sequencing technology used. In principle, any high-throughput sequencing technology²⁵ can be used for RNA-Seq, and the Illumina IG^{18–21,23,24}, Applied Biosystems SOLiD²² and Roche 454 Life Science^{26–28}

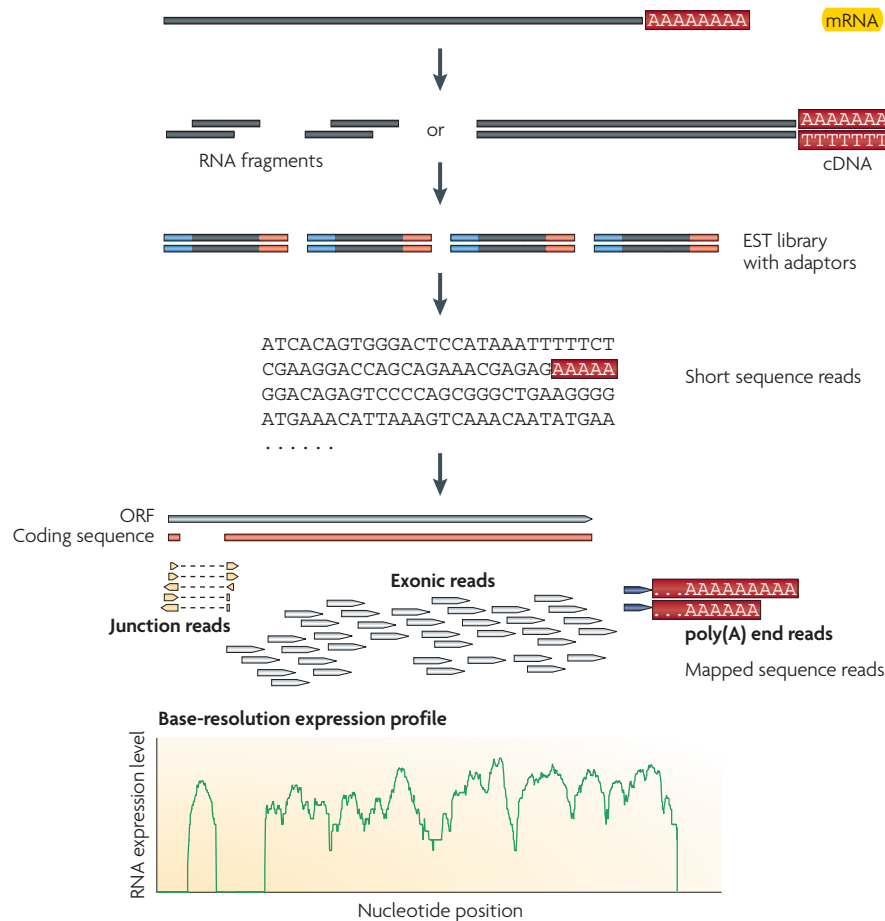


Figure 1 | A typical RNA-Seq experiment. Briefly, long RNAs are first converted into a library of cDNA fragments through either RNA fragmentation or DNA fragmentation (see main text). Sequencing adaptors (blue) are subsequently added to each cDNA fragment and a short sequence is obtained from each cDNA using high-throughput sequencing technology. The resulting sequence reads are aligned with the reference genome or transcriptome, and classified as three types: exonic reads, junction reads and poly(A) end-reads. These three types are used to generate a base-resolution expression profile for each gene, as illustrated at the bottom; a yeast ORF with one intron is shown.

systems have already been applied for this purpose. The *Helicos Biosciences* tSMS system has not yet been used for published RNA-Seq studies, but is also appropriate and has the added advantage of avoiding amplification of target cDNA. Following sequencing, the resulting reads are either aligned to a reference genome or reference transcripts, or assembled *de novo* without the genomic sequence to produce a genome-scale transcription map that consists of both the transcriptional structure and/or level of expression for each gene.

Although RNA-Seq is still a technology under active development, it offers several key advantages over existing technologies (TABLE 1). First, unlike hybridization-based approaches, RNA-Seq is not limited to detecting transcripts that correspond to existing genomic sequence. For

example, 454-based RNA-Seq has been used to sequence the transcriptome of the Glanville fritillary butterfly²⁷. This makes RNA-Seq particularly attractive for non-model organisms with genomic sequences that are yet to be determined. RNA-Seq can reveal the precise location of transcription boundaries, to a single-base resolution. Furthermore, 30-bp short reads from RNA-Seq give information about how two exons are connected, whereas longer reads or pair-end short reads should reveal connectivity between multiple exons. These factors make RNA-Seq useful for studying complex transcriptomes. In addition, RNA-Seq can also reveal sequence variations (for example, SNPs) in the transcribed regions^{22,24}.

A second advantage of RNA-Seq relative to DNA microarrays is that RNA-Seq has very low, if any, background

signal because DNA sequences can be unambiguously mapped to unique regions of the genome. RNA-Seq does not have an upper limit for quantification, which correlates with the number of sequences obtained. Consequently, it has a large dynamic range of expression levels over which transcripts can be detected: a greater than 9,000-fold range was estimated in a study that analysed 16 million mapped reads in *Saccharomyces cerevisiae*¹⁸, and a range spanning five orders of magnitude was estimated for 40 million mouse sequence reads²⁰. By contrast, DNA microarrays lack sensitivity for genes expressed either at low or very high levels and therefore have a much smaller dynamic range (one-hundredfold to a few-hundredfold) (FIG. 2). RNA-Seq has also been shown to be highly accurate for quantifying expression levels, as determined using quantitative PCR (qPCR)¹⁸ and spike-in RNA controls of known concentration²⁰. The results of RNA-Seq also show high levels of reproducibility, for both technical and biological replicates^{18,22}. Finally, because there are no cloning steps, and with the Helicos technology there is no amplification step, RNA-Seq requires less RNA sample.

Taking all of these advantages into account, RNA-Seq is the first sequencing-based method that allows the entire transcriptome to be surveyed in a very high-throughput and quantitative manner. This method offers both single-base resolution for annotation and 'digital' gene expression levels at the genome scale, often at a much lower cost than either tiling arrays or large-scale Sanger EST sequencing.

Challenges for RNA-Seq

Library construction. The ideal method for transcriptomics should be able to directly identify and quantify all RNAs, small or large. Although there are only a few steps in RNA-Seq (FIG. 1), it does involve several manipulation stages during the production of cDNA libraries, which can complicate its use in profiling all types of transcript.

Unlike small RNAs (microRNAs (miRNAs), Piwi-interacting RNAs (piRNAs), short interfering RNAs (siRNAs) and many others), which can be directly sequenced after adaptor ligation, larger RNA molecules must be fragmented into smaller pieces (200–500 bp) to be compatible with most deep-sequencing technologies. Common fragmentation methods include

Table 1 | **Advantages of RNA-Seq compared with other transcriptomics methods**

Technology	Tiling microarray	cDNA or EST sequencing	RNA-Seq
Technology specifications			
Principle	Hybridization	Sanger sequencing	High-throughput sequencing
Resolution	From several to 100 bp	Single base	Single base
Throughput	High	Low	High
Reliance on genomic sequence	Yes	No	In some cases
Background noise	High	Low	Low
Application			
Simultaneously map transcribed regions and gene expression	Yes	Limited for gene expression	Yes
Dynamic range to quantify gene expression level	Up to a few-hundredfold	Not practical	>8,000-fold
Ability to distinguish different isoforms	Limited	Yes	Yes
Ability to distinguish allelic expression	Limited	Yes	Yes
Practical issues			
Required amount of RNA	High	High	Low
Cost for mapping transcriptomes of large genomes	High	High	Relatively low

RNA fragmentation (RNA hydrolysis or nebulization) and cDNA fragmentation (DNase I treatment or sonication). Each of these methods creates a different bias in the outcome. For example, RNA fragmentation has little bias over the transcript body²⁰, but is depleted for transcript ends compared with other methods (FIG. 3). Conversely, cDNA fragmentation is usually strongly biased towards the identification of sequences from the 3' ends of transcripts, and thereby provides valuable information about the precise identity of these ends¹⁸ (FIG. 4).

Some manipulations during library construction also complicate the analysis of RNA-Seq results. For example, many short reads that are identical to each

other can be obtained from cDNA libraries that have been amplified. These could be a genuine reflection of abundant RNA species, or they could be PCR artefacts. One way to discriminate between these possibilities is to determine whether the same sequences are observed in different biological replicates.

Another key consideration concerning library construction is whether or not to prepare strand-specific libraries, as has been done in two studies^{21,22}. These libraries have the advantage of yielding information about the orientation of transcripts, which is valuable for transcriptome annotation, especially for regions with overlapping transcription from opposite directions^{2,19,29}; however, strand-specific libraries are

currently laborious to produce because they require many steps²² or direct RNA–RNA ligation²¹, which is inefficient. Moreover, it is essential to ensure that the antisense transcripts are not artefacts of reverse transcription³⁰. Because of these complications, most studies thus far have analysed cDNAs without strand information.

Bioinformatic challenges. Like other high-throughput sequencing technologies, RNA-Seq faces several informatics challenges, including the development of efficient methods to store, retrieve and process large amounts of data, which must be overcome to reduce errors in image analysis and base-calling and remove low-quality reads.

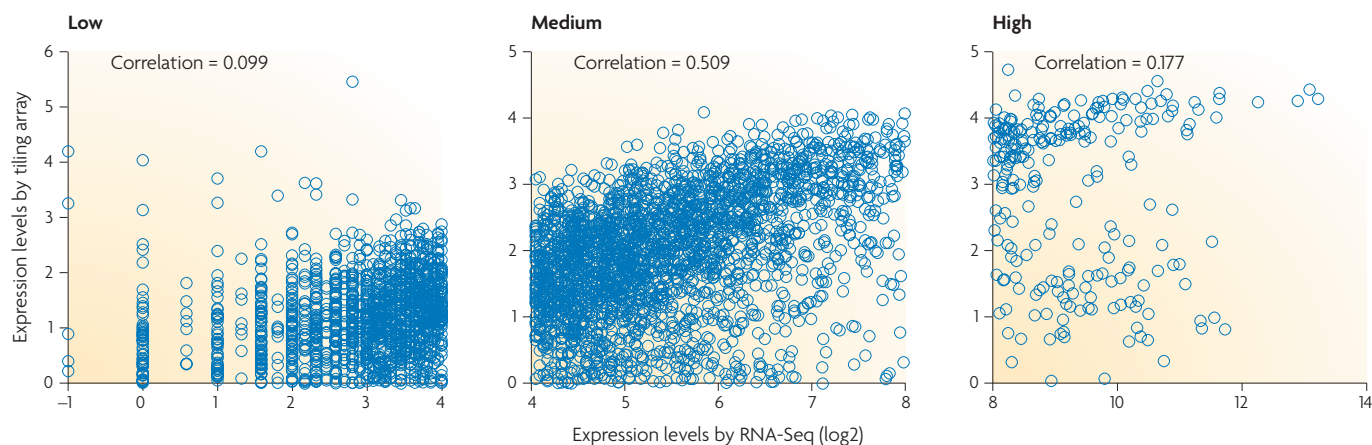


Figure 2 | **Quantifying expression levels: RNA-Seq and microarray compared.** Expression levels are shown, as measured by RNA-Seq and tiling arrays, for *Saccharomyces cerevisiae* cells grown in nutrient-rich media. The two methods agree fairly well for genes with

medium levels of expression (middle), but correlation is very low for genes with either low or high expression levels. The tiling array data used in this figure is taken from REF. 2, and the RNA-Seq data is taken from REF. 18.

Once high-quality reads have been obtained, the first task of data analysis is to map the short reads from RNA-Seq to the reference genome, or to assemble them into contigs before aligning them to the genomic sequence to reveal transcription structure. There are several programs for mapping reads to the genome, including ELAND, SOAP³¹, MAQ³² and RMAP³³ (information about these can be found at the [Illumina forum](#) and at [SEQanswers](#)). However, short transcriptomic reads also contain reads that span exon junctions or that contain poly(A) ends — these cannot be analysed in the same way. For genomes in which splicing is rare (for example, *S. cerevisiae*) special attention

only needs to be given to poly(A) tails and to a small number of exon–exon junctions. Poly(A) tails can be identified simply by the presence of multiple As or Ts at the end of some reads. Exon–exon junctions can be identified by the presence of a specific sequence context (the GT–AG dinucleotides that flank splice sites) and confirmed by the low expression of intronic sequences, which are removed during splicing. Transcriptome maps have been generated in this manner for *S. cerevisiae*¹⁸. For complex transcriptomes it is more difficult to map reads that span splice junctions, owing to the presence of extensive alternative splicing and trans-splicing. One partial solution is to compile

a junction library that contains all the known and predicted junction sequences and map reads to this library^{19,20}. A challenge for the future is to develop computationally simple methods to identify novel splicing events that take place between two distant sequences or between exons from two different genes.

For large transcriptomes, alignment is also complicated by the fact that a significant portion of sequence reads match multiple locations in the genome. One solution is to assign these multi-matched reads by proportionally assigning them based on the number of reads mapped to their neighbouring unique sequences^{20,22}. This method has been successful for low-copy repetitive sequences²⁰. Short reads that have high copy numbers (>100) and long stretches of repetitive regions present a greater challenge. Obtaining longer sequence reads, for example using 454 technology, should help alleviate the multi-matching problem. Alternatively, a paired-end sequencing strategy, in which short sequences are determined from both ends of a DNA fragment^{25,34,35}, extends the mapped fragment length to 200–500 bp and is expected to be useful in the future. Sequencing errors and polymorphisms can present mapping problems for all genomes, not just for repetitive DNA. Generally, single base differences are not problematic, because most mapping algorithms accommodate one or two base differences. However, resolving larger differences will require better reference genome annotation for polymorphisms and deeper sequencing coverage.

Coverage versus cost. Another important issue is sequence coverage, or the percentage of transcripts surveyed, which has implications for cost. Greater coverage requires more sequencing depth. To detect a rare transcript or variant, considerable depth is needed. In simple transcriptomes, such as yeast (both *S. pombe* and *S. cerevisiae*) for which there is no evidence of alternative splicing, 30 million 35-nucleotide reads from poly(A) mRNA libraries are sufficient to observe transcription from most (>90%) genes for cells grown under a single condition (that is, in nutrient-rich medium)¹⁸. This depth is probably more than sufficient for most purposes, as the number of expressed genes detected by RNA-Seq reaches 80% coverage at 4 million uniquely mapped reads, after which doubling the depth merely increases the coverage by 10% (FIG. 5). The remaining

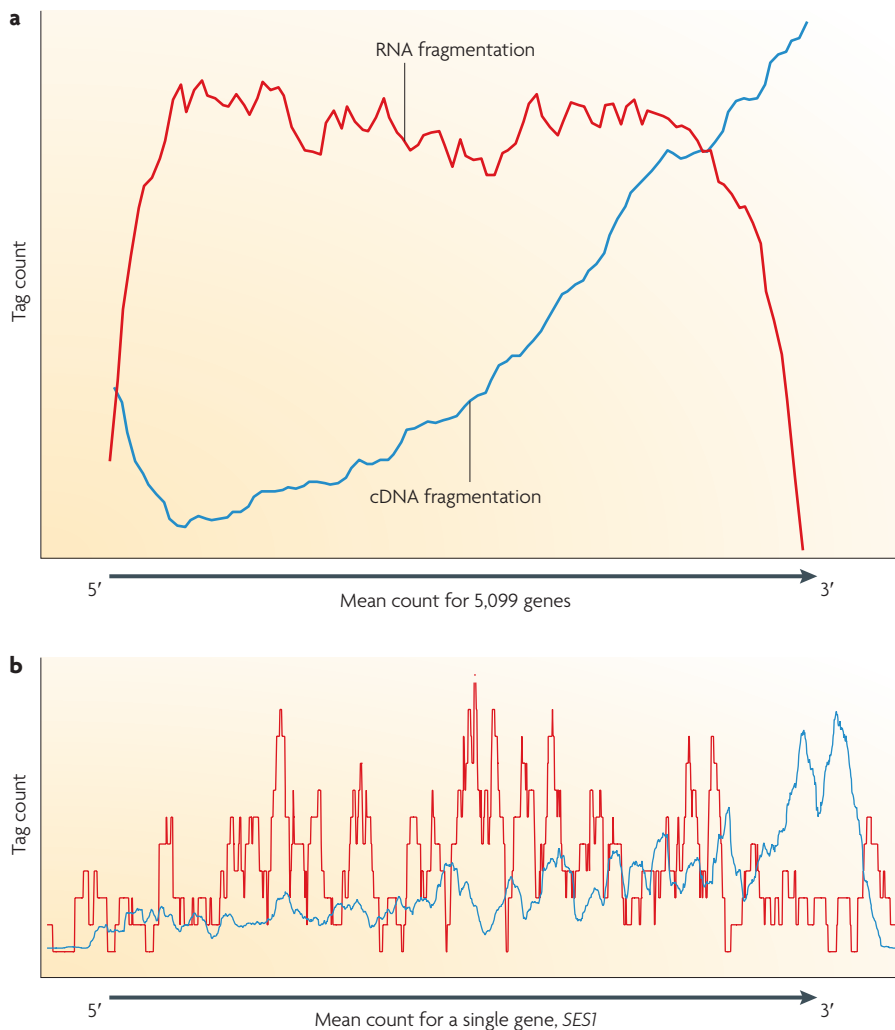


Figure 3 | DNA library preparation: RNA fragmentation and DNA fragmentation compared. **a** | Fragmentation of oligo-dT primed cDNA (blue line) is more biased towards the 3' end of the transcript. RNA fragmentation (red line) provides more even coverage along the gene body, but is relatively depleted for both the 5' and 3' ends. Note that the ratio between the maximum and minimum expression level (or the dynamic range) for microarrays is 44, for RNA-Seq it is 9,560. The tag count is the average sequencing coverage for 5,000 yeast ORFs¹⁸. **b** | A specific yeast gene, *SES1* (seryl-tRNA synthetase), is shown.

genes are presumably either not expressed under this condition (for example, sporulation genes¹⁸) or do not have poly(A) tails. Analyzing many different conditions can further increase the coverage; in *S. pombe* 122 million reads from six different growth conditions detected transcription from >99% of annotated genes¹⁹.

In general, the larger the genome, the more complex the transcriptome, the more sequencing depth is required for adequate coverage. Unlike genome-sequencing coverage, it is less straightforward to calculate the coverage of the transcriptome; this is because the true number and level of different transcript isoforms is not usually known and because transcription activity varies greatly across the genome. One study used the number of unique transcription start sites as a measure of coverage in mouse embryonic cells, and demonstrated that at 80 million reads, the number of start sites reached a plateau²² (FIG. 5b). However, this approach does not address transcriptome complexity in alternative splicing and transcription termination sites; presumably further sequencing can reveal additional variants.

New transcriptomic insights

Despite the challenges described above, the advantages of RNA-Seq have enabled us to generate an unprecedented global view of the transcriptome and its organization for a number of species and cell types. Before the advent of RNA-Seq, it was known that a much greater than expected fraction of the yeast, *Drosophila melanogaster* and human genomes are transcribed^{2,4,36}, and for yeast and humans a number of distinct isoforms have been found for many genes^{2,4}. However, the starts and ends of most transcripts and exons had not been precisely resolved and the extent of spliced heterogeneity remained poorly understood. RNA-Seq, with its high resolution and sensitivity has revealed many novel transcribed regions and splicing isoforms of known genes, and has mapped 5' and 3' boundaries for many genes.

Mapping gene and exon boundaries. The single-base resolution of RNA-Seq has the potential to revise many aspects of the existing gene annotation, including gene boundaries and introns for known genes as well as the identification of novel transcribed regions. 5' and 3' boundaries can be mapped to within 10–50 bases by a precipitous drop in signal. 3' boundaries

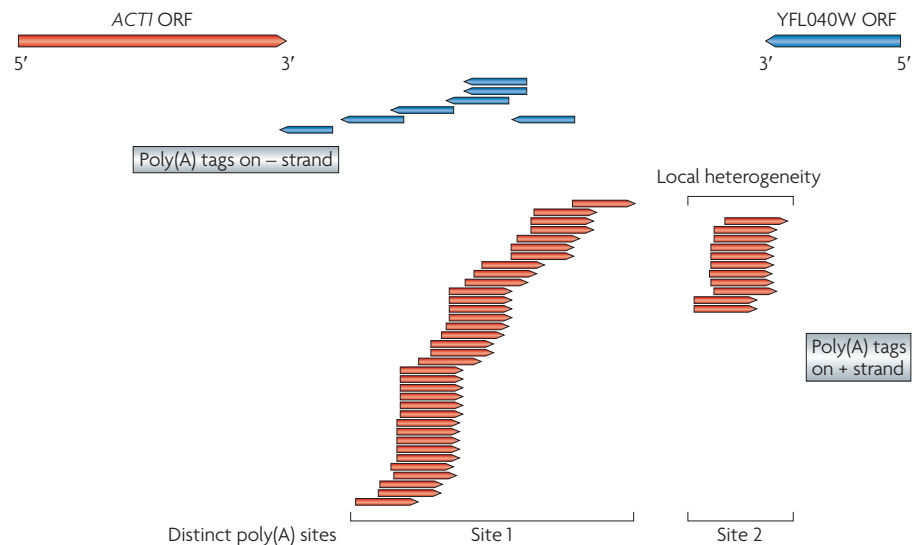


Figure 4 | Poly(A) tags from RNA-Seq. A region containing two overlapping transcripts (ACT1, from the actin gene, and YFL040W, an uncharacterized ORF) from the *Saccharomyces cerevisiae* genome is shown. Arrows point to transcription direction. The poly(A) tags from RNA-Seq experiments are shown below these transcripts, with arrows indicating transcription direction. The precise location of each locus identified by poly(A) tags reveals the heterogeneity in poly(A) sites, for example, ACT1 has two big clusters, both with a few bases of local heterogeneity. The transcription direction revealed by poly(A) tags also helps to resolve 3'-end overlapping transcribed regions¹⁸.

can be precisely mapped by searching for poly(A) tags, and introns can be mapped by searching for tags that span GT–AG splicing consensus sites. Using these methods the 5' and 3' boundaries of 80% and 85% of all annotated genes, respectively, were mapped in *S. cerevisiae*¹⁸. Similarly, in *S. pombe* many boundaries were defined by RNA-Seq data in combination with tiling array data¹⁹.

These two studies led to the discovery of many 5' and 3' UTRs that had not been analysed previously. In *S. cerevisiae*, extensive 3'-end heterogeneity was discovered at two levels: first, local heterogeneity exists in which a cluster of sites are involved, typically within a 10 bp window; second, there are distinct regions of poly(A) addition for 540 genes (FIG. 4). It is plausible that these different 3' ends confer distinct properties to the different mRNA isoforms, such as mRNA localization or degradation signals, which in turn might be responsible for unique biological functions^{18,19}. In addition to 3' heterogeneity, the list of upstream ORFs within the 5' UTRs of mRNAs (uORFs) was also greatly expanded from 17 to 340 (6% of yeast genes)¹⁸; uORFs regulate mRNA translation³⁷ or stability³⁸, so these sequences might make a previously underappreciated contribution to the regulatory sophistication of eukaryotic genomes. Interestingly,

many mRNAs with uORFs are transcription factors, suggesting that these regulators are themselves heavily regulated.

The mapping of transcript boundaries revealed several novel features of eukaryotic gene organization. Many yeast genes were found to overlap at their 3' ends¹⁸. Using relaxed criteria similar to those employed in a recent study¹⁸ we found that 808 pairs, approximately 25% of all yeast ORFs, overlap at their 3' ends¹⁸. Likewise, antisense expression is enriched in the 3' exons of mouse transcripts²². These features might confer interesting regulatory properties on the affected genes. For multicellular organisms, antisense transcription could modulate gene expression through the production of siRNAs or through dsRNA editing^{39,40}. For yeast, which seems to lack siRNA and dsRNA-editing functions, transcription from one gene might interfere with that from an overlapping gene, or coordinate gene expression through other mechanisms.

Extensive transcript complexity. RNA-Seq can be used to quantitatively examine splicing diversity by searching for reads that span known splice junctions as well as potential new ones. In humans, 31,618 known splicing events were confirmed (11% of all known splicing events) and 379 novel splicing events were discovered²⁴.

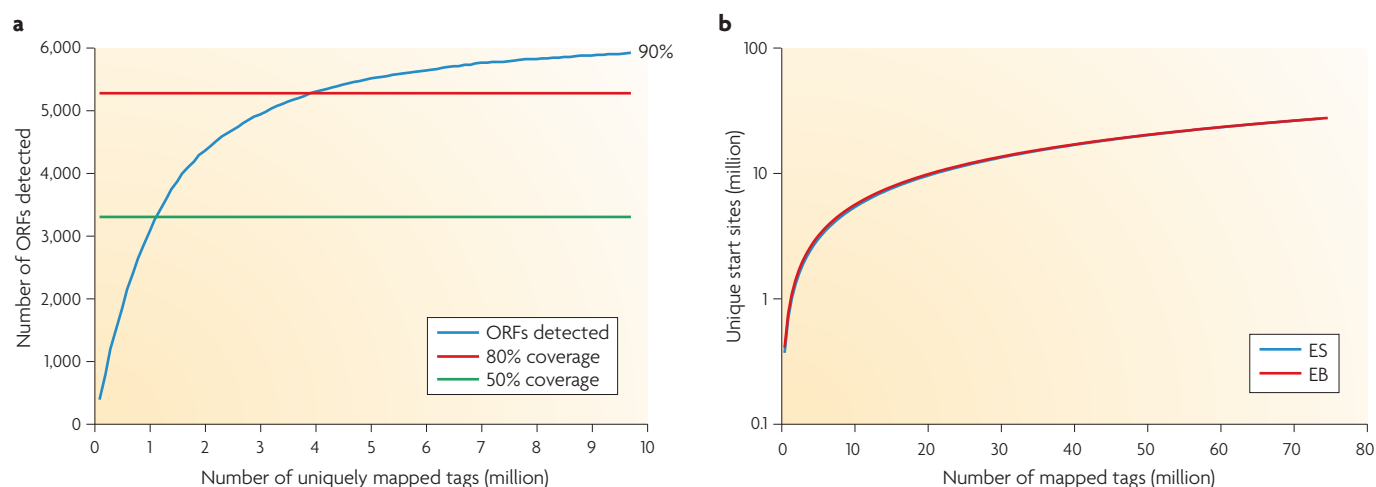


Figure 5 | Coverage versus depth. a | 80% of yeast genes were detected at 4 million uniquely mapped RNA-Seq reads, and coverage reaches a plateau afterwards despite the increasing sequencing depth. Expressed genes are defined as having at least four independent reads from a 50-bp window at the 3' end. Data is taken from REF. 18.

b | The number of unique start sites detected starts to reach a plateau when the depth of sequencing reaches 80 million in two mouse transcriptomes. ES, embryonic stem cells; EB, embryonic body. Figure is modified, with permission, from REF. 22 © (2008) Macmillan Publishers Ltd. All rights reserved.

Another study of human cells found 94,241 junctions, among which 4,096 were novel, and further demonstrated that the prevalent form of alternative splicing is exon skipping⁴¹. In mice, extensive alternative splicing was observed for 3,462 genes²⁰. In addition, 42 splicing events that join exons from multiple mouse genes were detected²².

Novel transcription. Previous studies using transposon tagging and tiling microarrays have suggested that in the genomes of yeast, *D. melanogaster* and humans, there are many novel transcribed regions

represented in poly(A)+ RNA^{2,36,42,43}. However, the accuracy of the tiling array results is uncertain owing to concerns about cross-hybridization (see below). RNA-Seq, which does not suffer from problems with background noise, has confirmed that at least 75% and perhaps greater than 90% of the *S. cerevisiae* and *S. pombe* genomes are expressed^{18,19}. In addition, results from RNA-Seq suggest the existence of a large number of novel transcribed regions in every genome surveyed, including the *A. thaliana*²¹, mouse^{20,22}, human²⁴, *S. cerevisiae*¹⁸ and *S. pombe*¹⁹ genomes. 487 and 453 novel transcripts have been discovered in

S. cerevisiae and *S. pombe*, respectively^{18,19}; for *S. cerevisiae* half of these were not identified using microarrays. Many of these novel transcribed regions in yeast do not seem to encode any protein, and their functions remain to be determined. The current sequencing depth is not sufficient to define the boundaries of novel transcript units in mammals; however, 30–40% of reads map to unannotated regions^{20,22,24}. These novel transcribed regions, combined with many undiscovered novel splicing variants, suggest that there is considerably more transcript complexity than previously appreciated.

Glossary

Cap analysis of gene expression

(CAGE). Similar to SAGE, except that 5'-end information of the transcript is analysed instead of 3'-end information.

Contigs

A group of sequences representing overlapping regions from a genome or transcriptome.

dsRNA editing

Site-specific modification of a pre-mRNA by dsRNA-specific enzymes that leads to the production of variant mRNA from the same gene.

Genomic tiling microarray

A DNA microarray that uses a set of overlapping oligonucleotide probes that represent a subset of or the whole genome at very high resolution.

Massively parallel signature sequencing

(MPSS). A gene expression quantification method that determines 17–20-bp 'signatures' from the ends of a cDNA molecule using multiple cycles of enzymatic cleavage and ligation.

MicroRNA

(miRNA). Small RNA molecules that are processed from small hairpin RNA (shRNA) precursors that are produced from miRNA genes. miRNAs are 21–23 nucleotides in length and through the RNA-induced silencing complex they target and silence mRNAs containing imperfectly complementary sequence.

Piwi-interacting RNAs

(piRNA). Small RNA species that are processed from single-stranded precursor RNAs. They are 25–35 nucleotides in length and form complexes with the piwi protein. piRNAs are probably involved in transposon silencing and stem-cell function.

Quantitative PCR

(qPCR). An application of PCR to determine the quantity of DNA or RNA in a sample. The measurements are often made in real time and the method is also called real-time PCR.

Sequencing depth

The total number of all the sequences reads or base pairs represented in a single sequencing experiment or series of experiments.

Serial analysis of gene expression

(SAGE). A method that uses short ~14–20-bp sequence tags from the 3' ends of transcripts to measure gene expression levels.

Short interfering RNA

(siRNA). RNA molecules that are 21–23 nucleotides long and that are processed from long double-stranded RNAs; they are functional components of the RNAi-induced silencing complex. siRNAs typically target and silence mRNAs by binding perfectly complementary sequences in the mRNA and causing their degradation and/or translation inhibition.

Spike-in RNA

A few species of RNA with known sequence and quantity that are added as internal controls in RNA-Seq experiments.

Defining transcription level

As RNA-Seq is quantitative, it can be used to determine RNA expression levels more accurately than microarrays. In principle, it is possible to determine the absolute quantity of every molecule in a cell population, and directly compare results between experiments. Several methods have been used for quantification. For RNA fragmentation followed by cDNA synthesis, which gives more uniform coverage of each exon, gene expression levels can be deduced from the total number of reads that fall into the exons of a gene, normalized by the length of exons that can be uniquely mapped²⁴; for 3'-biased methods, read counts from a window near the 3' end are used¹⁸. Gene expression levels determined by these methods closely correlate with qPCR and RNA spike-in controls.

One particularly powerful advantage of RNA-Seq is that it can capture transcriptome dynamics across different tissues or conditions without sophisticated normalization of data sets^{19,20,22}. RNA-Seq has been used to accurately monitor gene expression during yeast vegetative growth¹⁸, yeast meiosis¹⁹ and mouse embryonic stem-cell differentiation²², to track gene expression changes during development, and to provide a 'digital measurement' of gene expression difference between different tissues²⁰. Because of these advantages, RNA-Seq will undoubtedly be valuable for understanding transcriptomic dynamics during development and normal physiological changes, and in the analysis of biomedical samples, where it will allow robust comparison between diseased and normal tissues, as well as the subclassification of disease states.

Future directions

Although RNA-Seq is still in the early stages of use, it has clear advantages over previously developed transcriptomic methods. **The next big challenge for RNA-Seq is to target more complex transcriptomes to identify and track the expression changes of rare RNA isoforms from all genes.** Technologies that will advance achievement of this goal are pair-end sequencing, strand-specific sequencing and the use of longer reads to increase coverage and depth. As the cost of sequencing continues to fall, RNA-Seq is expected to replace microarrays for many applications that involve determining the structure and dynamics of the transcriptome.

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