# RNA-Seq methods for transcriptome analysis



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Deep sequencing has been revolutionizing biology and medicine in recent years, providing single base-level precision for our understanding of nucleic acid sequences in high throughput fashion. Sequencing of RNA, or RNA-Seq, is now a common method to analyze gene expression and to uncover novel RNA species. Aspects of RNA biogenesis and metabolism can be interrogated with specialized methods for cDNA library preparation. In this study, we review current RNA-Seq methods for general analysis of gene expression and several specific applications, including isoform and gene fusion detection, digital gene expression profiling, targeted sequencing and single-cell analysis. In addition, we discuss approaches to examine aspects of RNA in the cell, technical challenges of existing RNA-Seq methods, and future directions. © 2016 Wiley Periodicals, Inc.

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### INTRODUCTION

RNA molecules are essential components of all living cells. Understanding the identity and abundance of each RNA molecule in a given cell under a specific condition is the ultimate goal of RNA research. Much of what we know about RNA comes from studies using biochemical methods, where a small number of specific molecules are analyzed. High-throughput approaches that enable interrogation of RNA sequences on a large scale emerged in the early 1990s. The expressed sequence tag (EST) method developed by Adams et al. examines gene expression by partially sequencing complementary DNA (cDNA) clones, revealing both the sequence and the abundance of corresponding RNAs. EST data played a pivotal role in identification of new genes in genomes in the 1990s. However, the high sequencing cost of the method limited its use in expression analysis, and the data is largely believed to be semi-quantitative. The Serial Analysis of Gene

The first decade of this millennium witnessed the advent of massive parallel sequencing, also known as deep sequencing or Next Generation Sequencing (NGS). Lauded as revolutionary in biology and medicine for its ability to acquire an unprecedented amount of data in a short time, deep sequencing quickly transformed RNA research.

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Expression (SAGE) method developed by Velculescu et al. significantly cut down the cost of expression analysis on a per gene basis,<sup>2</sup> thanks to sequencing only a short tag region per cDNA (15 bp for the short SAGE method and 21 bp for the long SAGE method). However, the emergence of DNA microarray technology in the mid-1990s superseded EST and SAGE methods for gene expression analysis, largely due to its much better affordability for large scale studies.<sup>3,4</sup> DNA microarray analysis of gene expression is based on hybridization of fluorescently labeled targets that are derived from transcripts to probes that are attached to a solid surface through printing or in situ synthesis. However, while the method enables interrogation of transcripts genome-wide, the requirement that a priori sequence information or reference genomes/transcriptomes be available for designing the microarray probes limited the development and application of this technology in discovery applications. In addition, cross-hybridization and background signals often lead to low specificity or low sensitivity for some genes.

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RNA-Seq is now the method of choice to study gene expression and identify novel RNA species. Compared to DNA microarray-based methods, RNA-Seq offers less background noise and a greater dynamic range for detection. Most importantly, RNA-Seq directly reveals sequence identity, crucial for analysis of unknown genes and novel transcript isoforms. Several different technologies have been developed for RNA-Seq. Here we review general aspects of RNA-Seq and applications of RNA-Seq to study specific problems. We discuss issues and remedies related to bias and sensitivity in these methods, two paramount concerns in an RNA-Seq experiment. We also discuss specialized methods that investigate aspects of RNA biogenesis and metabolism.

### GENERAL ASPECTS OF RNA-Seq

While direct sequencing of RNA molecules is possible, 10 most RNA-Seq experiments are carried out on instruments that sequence DNA molecules due to the technical maturity of commercial instruments designed for DNA-based sequencing. Therefore, cDNA library preparation from RNA is a required step for RNA-Seq. Each cDNA in an RNA-Seg library is composed of a cDNA insert of certain size flanked by adapter sequences, as required for amplification and sequencing on a specific platform. The cDNA library preparation method varies depending on the RNA species under investigation, which can differ in size, sequence, structural features and abundance. Major considerations include (1) how to capture RNA molecules of interest; (2) how to convert RNA to double-stranded cDNAs with defined size ranges; and (3) how to place adapter sequences on the cDNA ends for amplification and sequencing. These are discussed in the following sections.

### Selection of Poly(A) + Transcripts

Sequencing of polyadenylated RNA is perhaps the most common application of RNA-Seq. In eukaryotic organisms, most protein-coding RNAs (mRNAs) and many long noncoding RNAs (lncRNAs) (>200 nt) contain a poly(A) tail. The poly(A) tail provides technical convenience for enrichment of poly(A) + RNAs from total cellular RNA, in which they account for approximately 1–5% of the pool. Poly(A) + RNA selection can be carried out with magnetic or cellulose beads coated with oligo-dT molecules. Alternatively, polyadenylated RNAs can be selected using oligo-dT priming for reverse transcription (RT). While efficiently incorporating both poly(A) selection

and RT in one step, oligo-dT priming-based methods can exhibit 3' bias, resulting in sequencing reads enriched for the 3' portion of the transcript. In addition, oligo-dT can frequently prime at internal A-rich sequences of transcripts, a phenomenon called internal poly(A) priming,<sup>7,11</sup> leading to biased RT. Therefore, poly(A) purification is a preferred method to select poly(A) + RNA unless a very low amount of RNA is available (see next).

### rRNA Depletion

Non-polyadenylated RNAs, such as prokaryotic mRNAs, fragmented mRNAs from formalin-fixed, paraffin-embedded (FFPE) samples, and poly(A)-transcripts in eukaryotic cells are often the subject of investigation. A major issue in sequencing these RNAs is how to eliminate ribosomal RNAs (rRNAs), which are the most abundant RNA species in the cell but of little interest in most studies. Several approaches have been developed to deplete them from the RNA pool.

One approach to eliminate rRNAs is based on sequence-specific probes that can hybridize to rRNAs. Unwanted rRNAs or their cDNAs are hybridized with biotinylated DNA or locked nucleic acid (LNA) probes, followed by depletion with streptavidin beads. Alternatively, rRNAs are targeted by antisense DNA oligos and digested by RNase H, a method also known as probe-directed degradation (PDD). While this approach is less laborious than hybridization, it requires continuous coverage of rRNAs and unique probe sets designed for different species. A noncontinuous sequence-based method was recently developed which has addressed some of these issues. 12,13 In this method, all cDNAs, including those of rRNAs and other RNAs, are circularized, and are hybridized to rRNA probes. The hybridized sequences are then digested by duplexspecific nuclease (DSN), making them unusable for amplification. However, this approach requires high input amounts of total RNA, which can be challenging when dealing with clinical samples.

Another approach for rRNA reduction uses specific, not-so-random (NSR) primers which bind to the RNA molecules of interest during RT, thus avoiding rRNAs. This method, commercialized by NuGEN under the name Ovation RNA-Seq, uses hexamer or heptamer primers whose sequences are absent from rRNAs. Similar to this approach, one study used 44 heptamers to avoid both rRNAs and highly-expressed transcripts. A recent report used only 40 primers for RT instead of 700 NSR primers commonly used in other studies. A key advantage of

this approach is that NSR primers work well with partially degraded RNA and low-input samples. However, like other sequence targeting methods, this approach suffers from off-target priming, and is species dependent. Nevertheless, NSR primers are frequently used in prokaryotic species, for which poly(A) purification is not an option.

In addition to the sequence-based approaches mentioned above, some methods take advantage of certain features of rRNAs for their elimination. The C<sub>0</sub>T-hybridization method is based on heat denaturation, re-annealing and selective degradation by DSN. Double-stranded cDNAs originated from abundant sequences are preferentially degraded because of their more rapid annealing kinetics compared to less abundant ones. Selective degradation has also been achieved by using the enzyme terminator 5'-phosphate-dependent exonuclease (TEX), which recognizes RNA molecules with 5'-monophosphate, as with rRNAs and tRNAs. 19

In summary, the selection of an approach for enriching RNA transcripts of interest for sequencing depends on the goal of the experiment and many technical factors. Several studies have compared protocols for removal of rRNA by depletion- and priming-based methods.<sup>7,20–23</sup> In eukaryotic cells, oligo-dT bead-based purification of poly(A) + RNA is the method of choice for most applications, because of its ease of use and relatively low cost. For low-input samples, however, oligo-dT priming generally offers better results. Both poly(A) selection methods can effectively address intron sequence contamination. If an RNA sample is partially degraded or the user is interested in noncoding RNAs, depletion of rRNAs by PDD or NSR-priming is typically necessary. PDD works better for highinput samples whereas NSR-priming is primarily used for lower inputs of RNA.

### Fragmentation

After poly(A) + selection or rRNA depletion, RNA samples are typically subject to RNA fragmentation to a certain size range before RT. This is necessary because of the size limitation of most current sequencing platforms, e.g., <600 bp on Illumina sequencers. RNAs can be fragmented with alkaline solutions, solutions with divalent cations, such Mg++, Zn++, or enzymes, such RNase III. Fragmentation with alkaline solutions or divalent cations is typically carried out at an elevated temperature, such as 70°C, to mitigate the effect of RNA structure on fragmentation. Nevertheless, RNA fragmentation by chemical fragmentation is not completely random. Similarly,

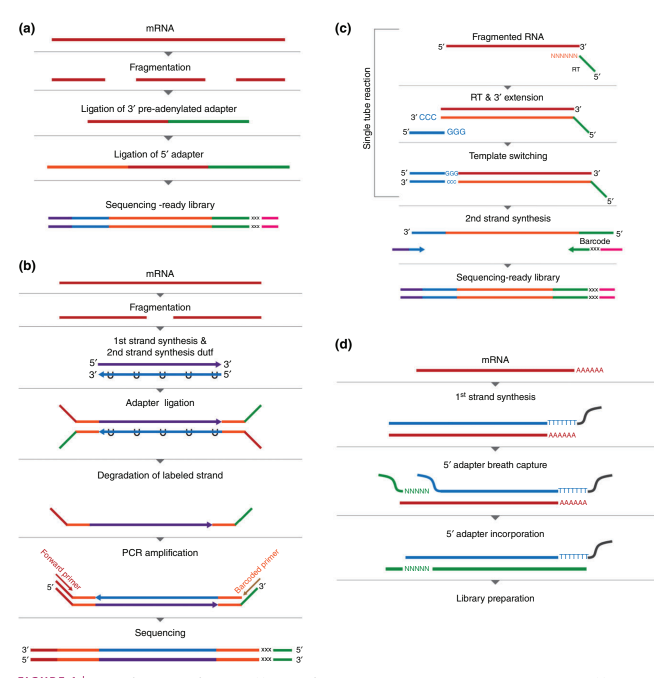
the RNase III-based method can also introduce bias because of the enzyme's preference for double-stranded RNA sequences.<sup>24</sup> Thus, uneven fragmentation of RNA can be a source of bias, leading to differential representation of specific regions of RNA.

Alternatively, intact RNAs can be reverse transcribed, and full-length cDNA can be fragmented. A traditional method to fragment cDNA requires the use of acoustic shearing, which is less amenable to automation than RNA fragmentation. Alternatively, full-length double-stranded cDNAs can be fragmented by DNases. Recent development in using transposon-based, so-called tagmentation method has made it simple to fragment cDNA and add adapter sequences at the same time.<sup>25</sup> In this method, an active variant of the Tn5 transposase mediates the fragmentation of double-stranded DNA and ligates adapter oligonucleotides at both ends in a quick reaction (~5 min). However, it is notable that Tn5 and other enzyme-based cDNA fragmentation methods require a precise enzyme:DNA ratio, making method optimization less straightforward than RNA fragmentation. Consequently, fragmenting RNA is currently still the most frequently used approach in RNA-Seq library preparation.

### Adapters and Directionality

In a standard RNA-Seq library protocol, cDNAs of a desired size generated from RT of fragmented RNAs with random hexamer primers or from fragmented full-length cDNAs are ligated to DNA adapters before amplification and sequencing. While simple, this approach loses the information about which DNA strand corresponds to the sense strand of RNA. Lack of strand specificity would make it difficult to identify antisense and novel RNA species and cause inaccurate measurement of sense RNA expression. Several methods have been developed to capture the directionality of RNA in cDNA libraries.<sup>26</sup>

The first approach involves attaching different adapters directly to the 5' and 3' ends of the RNA molecule (Figure 1(a)). Originally designed for small RNA-Seq,<sup>27</sup> this method begins with removal of 3' phosphate group from fragmented RNA and addition of a 5' phosphate group. This is followed by sequential ligations of a 5' adenylated 3' adapter using a truncated RNA ligase II and a 5' adapter ligation using RNA ligase I. The sequence difference between 5' and 3' adapters preserves RNA strandedness. While simple to implement, this approach suffers from substantial biases due to the influence of both 5' and 3' end sequences on the ligation steps. This issue, however, has recently been mitigated



**FIGURE 1** | Methods for strand-specific RNA-Seq. (a) Ligation of the 3' preadenylated and 5' adapters. "xxx" indicates barcode. (b) Labeling of the second strand with dUTP, followed by enzymatic degradation. (c) The Peregrine method involves template-switch attachment of the 3' adapter. (d) BrAD-Seq captures the 3' adapter by taking advantage of terminal breathing of double-stranded DNA.

significantly by using random nucleotides at the ligation end of each adapter. <sup>28,29</sup>

A second approach incorporates dUTP into the second strand of cDNA (Figure 1(b)). The labeled strand can be degraded before PCR amplification with uracil DNA glycosylase (UDG), an enzyme that cleaves the uracil base in dUTP-containing DNA. In addition, the U-containing strand is a very poor template for thermostable polymerases, such as the

Phusion polymerase, making it essentially not amplifiable. As such, only the first strand cDNA with defined adapter sequences is amplified, conferring directional information to the sequencing reads. A systematic comparison of different protocols for strand-specific RNA-Seq indicated that the dUTP-based method was the most effective in terms of evenness of coverage. Indeed, this method is currently the most frequently used in commercial

directional RNA-Seq library preparation protocols. However, since it requires extra enzymatic and purification steps that are laborious and can cause material loss, it is generally not suitable for low-input samples.

There have been several attempts to develop new methods based on adding adapters to fragmented RNA. One of these methods, Pedegrine, uses template-switching after priming with a randomhexamer that contains a small tag (Figure 1(c)).<sup>31</sup> Another method, BrAD-Seq, takes advantages of temporary strand separation in double-stranded DNA, called 'breathing,' to introduce a tag sequence (Figure 1(d)).<sup>32</sup> A third example is sequential ligation of tags to RNA that has been fragmented to 200 nt, preserving even small RNAs. 33,34 A DSN-mediated normalization step is used to remove rRNA products. A similar protocol for stranded RNA-Seq libraries from all RNA species has also been reported, 35 where an oligo containing a tag sequence for RT is ligated to the 3' end of RNA. A second tag is introduced by the RT primer. First strand synthesis is then followed by circularization of cDNA by DNA ligase and direct amplification of the library using the two tags.

### Amplification and Molecular Labels

Due to the detection limit of most sequencers, cDNA libraries need to be amplified by PCR before sequencing. While only a small number of amplification cycles (8-12) are used during PCR, variations in cDNA size and composition can result in uneven amplification. Amplification of some cDNAs plateau while others continue to amplify exponentially. To correct for PCR amplification bias, methods that eliminate PCR duplicates from sequencing results have been introduced. In one method, under the assumption of random RNA fragmentation, final sequencing reads having the same start and stop coordinates are considered as PCR duplicates and are merged. 36,37 However, a drawback of this approach is elimination of reads derived from frequently fragmented regions due to not-so-stochastic fragmentation.<sup>38</sup> Another method is to use molecular labels, also known as unique molecular identifiers (UMIs), to distinguish PCR products. 39-41 Molecular labels are typically introduced within the adapter sequence, prior to PCR amplification. In a modified protocol for making cDNAs from single cells, molecular labels were introduced by the Tn5 transposase during fragmentation of double-stranded, amplified cDNA<sup>42</sup>. However, in some applications, such as digital counting of targeted RNAs, molecular labels are added during RT. 40,43,44 Molecular labels differ in size (number of bases) and complexity. In principle, they

comprise either defined sequences or random nucleotides. Defined sequences, chosen for their even distribution in final libraries, are more technically challenging to make because of sequence selection and manufacturing complexity. By contrast, random sequences, while easy to implement, give high variability among molecular labels. Molecular labeling is particularly valuable in situations where input RNA is scarce and a large number of PCR cycles is required for sequencing, such as single-cell RNA-Seq (see next). 45

# RNA-Seq METHODS FOR SPECIFIC GOALS

## Tag-Based Methods for Gene Expression Profiling

### DGE-Seq

Digital gene expression (DGE)-Seq, or Tag-Seq, is a deep sequencing method derived from SAGE. 46,47 As in SAGE, the method involves attachment of mRNA to beads via the poly(A) tail, first and second strand cDNA syntheses on the beads, and digestion of double-stranded cDNA with a frequent cutting restriction enzyme. The remaining 3' fragment attached to the beads is then ligated to its 5' end adapter with a recognition site for another restriction enzyme, called tagging enzyme. The tagging enzyme cleaves the cDNA and generates a short 21 bp tag, which is then ligated to a second adapter at its 3' end. The cDNA is then amplified by PCR, followed by sequencing. Because only a short tag is sequenced from the whole transcript, DGE-Seq is more economical than traditional RNA-Seg for a given depth of sequencing and can provide a higher dynamic range of detection when the same number of reads is generated. By design, DGE-Seq preserves RNA strandedness. This method has been commercialized by several companies and is useful especially when simple gene expression profiling is the goal.<sup>48</sup> It is also the method of choice when the complete genome or transcriptome is not available for full alignment of RNA-Seq reads.

### 3' End Sequencing

A number of methods specifically sequence the 3' end region of transcripts. Most of these methods were first developed to interrogate alterative cleavage and polyadenylation sites, a widespread phenomenon in all eukaryotes. As in DGE-Seq, the data from these methods can also be used to study gene expression. Some of these methods use oligo(dT) to prime RT or sequencing, such as PAS-Seq, polyA-Seq, 13'T-

fill,<sup>52</sup> and so on. One concern with this approach is that internal poly(A) priming can generate a high frequency of truncated cDNA.11 Other methods, such as 3P-Seq<sup>53</sup> and 3'READS,<sup>54</sup> use RNA-based ligation to capture the 3' end fragments. While these methods successfully address the internal priming issue, sequence preference of RNA ligases can introduce bias. A more detailed review of these methods can be found in Ref 55. Also notable are methods that can examine both the 3' end region and the poly(A) tail length at the same time. TAIL-Seq adds an adapter to the 3' end of the poly(A) tail and carries out sequencing from both ends of the insert (paired-end sequencing) to reveal both the length of poly(A) tail and the sequence near the poly(A) site.<sup>56</sup> A related method, PAT-Seq, uses Klenow polymerase to add an adapter sequence to the 3' end of the poly(A) tail, and uses single-read sequencing to obtain the sequence near the poly(A) site, as well as part of or the whole poly(A) tail.<sup>57</sup>

In essence, DGE-Seq and 3' end sequencing are tag-based approaches that use one fragment to represent a transcript. While efficient for gene expression analysis, they can have higher variability than 'shotgun' style RNA-Seq, where one transcript is represented by multiple fragments. Biases from fragmentation, adapter ligation and PCR can make tag-based data more prone to batch effects.

### Sequencing to Reveal Alternative Splicing and Gene Fusion

Almost all multi-exon genes display alternative splicing (AS).<sup>58</sup> AS plays an important role in regulation of cellular processes, and aberrations of the process are associated with many human diseases.<sup>59</sup> Some RNA-Seq reads cover exon-exon junctions, providing direct evidence of AS. In addition, reads mapped to internal exonic regions can be used to predict the AS pattern using statistical inference methods.<sup>60</sup> A more direct approach to examine AS is to sequence the exon-exon junction region directly. Using oligo pairs targeted to specific exon-exon junction sequences, the Fu lab developed RASL-Seq,<sup>61</sup> which provides analysis of specific splice junction regions. However, prior knowledge of the exon-exon junction sequences is required for the oligo design.

Similar to splicing, gene fusion events can place two noncontinuous genomic regions together in a single transcript. Created by chromosomal rearrangements, gene fusions are present in approximately 20% of cancer. <sup>62</sup> Fusion events can be detected using RNA-Seq data along with specific bioinformatic methods. <sup>63–65</sup> Detection of a fusion event is typically

revealed by reads containing fusion junctions or by differences in expression between the 5' and 3' ends of genes that are fused. Regular RNA-Seq methods are typically not sufficiently sensitive to detect fusion junctions. Several methods have been developed, including (1) enrichment of RNA-Seq reads for genes of interest, (2) exon capture, and (3) amplicon sequencing. In a recent study, exon capture of 467 cancer-related genes was successfully employed for the detection of their fusion events.66 Amplicon targeting requires primer design at the 5' and 3' ends of a transcript, and allows quantitative analysis of fusion events. This strategy was recently employed for the detection of ALK fusion events in lung cancer samples.<sup>67</sup> Two similar amplicon-based methods have been developed and commercialized by NuGene and ArcherDX, 68,69 where expected fusion events were examined by two sequence-specific primers together with a common primer that targets adapter sequence.

The ultimate solution to unravel the complexity of alternatively splicing and gene fusion isoforms is to sequence each transcript from the beginning to the end. Two strategies have been established to this end. Single-molecule real-time sequencing (SMRT) on the PacBio sequencing platform offers long reads up to 5 kb.<sup>70</sup> However, this method is costly, and has a high error-rate and low multiplexing capacity. Hybrid sequencing methods have been introduced which combine SMRT data with short, standard RNA-Seq reads. 71-73 A second approach, named synthetic long-read-RNA sequencing (SLR-RNA-Seq), is based on the Illumina MOLECULO system.<sup>74</sup> With SLR RNA-Seq, an RNA substrate is diluted to no more than 1000 transcripts per well so that the probability that two transcripts from the same gene are in the same well is very low. As such, reads for a gene from each well can be assembled to cover the entire length of a single transcript of the gene. After fragmentation, barcoding, library preparation, and sequencing on the Illumina MOLECULO platform, the structures of the original transcripts in the pools can be delineated. Compared to PacBio sequencing, SLR-RNA-Seq was found to deliver longer transcripts and a greater number of detected isoforms.<sup>74</sup> These methods are particularly valuable for analysis of AS and aberrant fusion isoforms.

### Targeted RNA-Seq

Selection a specific set of transcripts for sequencing is often desirable when a defined group of genes is of interest. Lowly expressed genes that cannot be readily analyzed using whole transcriptome sequencing can also be detected using targeted RNA-Seq

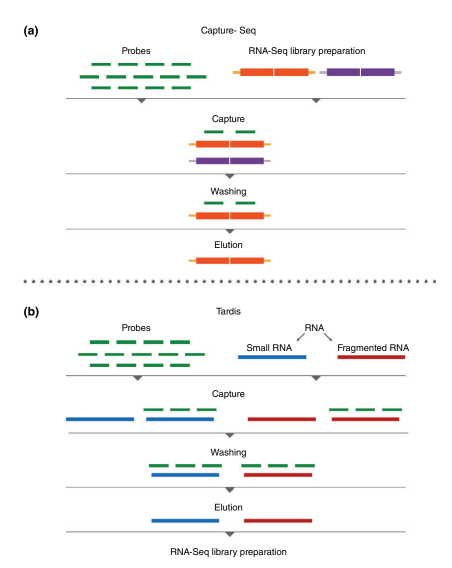
methods. Two general approaches have been used, namely, target capture and amplicon sequencing. The target capture approach involves selection of specific genes using a set of biotinylated probes which bind cDNA, <sup>66,75,76</sup> or RNA<sup>77</sup> (Figure 2). By contrast, amplicon sequencing employs gene-specific primers for the amplification of cDNA targets (Figure 3). Approaches differ in the amplicon design, including two specific primers after cDNA synthesis by template switch, <sup>78</sup> nested PCR with one specific primer and one common primer for adapter, <sup>69</sup> and specific targeting primers in combination with a primer for poly(A) tail priming. <sup>45</sup>

The target capture approach has been shown to provide greater complexity and uniformity than the

amplicon-based approach in a recent whole exome study.<sup>79</sup> However, target capture methods are more costly. Therefore, for studies that do not involve complex analysis, amplicon-based methods are preferred, as exemplified by analysis of T-cell and B-cell receptor repertoires.<sup>78,80,81</sup>

### Single-Cell RNA-Seq

RNA-Seq of cell populations gives rise to expression profiles that are averaged across cells. However, a cell bulk often contains different types or subtypes which are impossible to dissect using population-based analysis. In addition, the co-expression patterns between genes in a cell are lost when aggregating cells. Thus,



**FIGURE 2** | Targeted RNA-Seq by target capture. (a) The Capture-Seq method is based on capture of regions of interest by hybridization of RNA-Seq libraries to DNA oligonucleotide probes. (b) TARDIS is based on hybridization of input RNA to DNA oligonucleotide probes. The enrichment step is followed by the construction of a directional RNA-Seq library by ligation of 3' and 5' adapters.

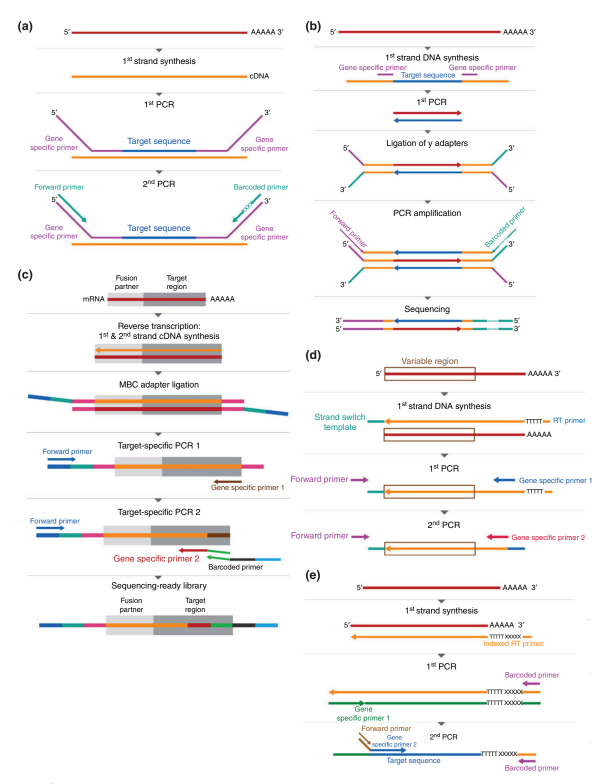


FIGURE 3 | Targeted RNA-Seq by amplicon sequencing. (a) PCR method using gene specific primers with overhangs containing sequences for common primers. (b) PCR methods using a pair of gene-specific primers followed by ligation of adapters with sequences necessary for sequencing. (c) Archer methods for detection of gene fusion. Sequence for a common primer is introduced by adapter ligation and is followed by nested PCR with gene-specific primers. MBC, Molecular Barcoded. (d) Detection of the TCR variable region. The sequence of a common primer is introduced during reverse transcription, and RT is followed by nested PCR with gene-specific primers. (e) Digital encoding of targeted mRNAs. Reverse transcription will introduce molecular indexes and sequences for common primers. Nested PCR follows, where gene-specific primers capture targeted sequences.

understanding gene expression at the single-cell level is important for obtaining the full picture of gene regulation in cells. Major challenges in single-cell analysis include isolation of single cells, sensitive methods to prepare cDNA libraries with very low inputs of RNA, and computational methods tailored for single-cell analysis. Several recent reviews have summarized these technologies and issues related to single-cell research. 82–86 Here, we focus on the library preparation aspect for single-cell RNA-Seq (Figure 4).

A single mammalian cell contains approximately 5–15 pg of RNA. However, the RNA-Seq methods described above are generally not suitable for sequencing RNA with less than 1 ng. Thus special RNA/DNA amplification or enhanced efficiency for sample processing is needed for single-cell RNAseq. Some methods, such as CEL-Seq and MARS-Seq, introduce a T7 promoter sequence with oligo(dT) during RT, which enables linear amplification of input RNA by in vitro transcription. 87,88 Second strand cDNA synthesis has been enhanced by template-switching during RT (SMART-Seq)<sup>89</sup> and poly(A) tailing of cDNA.<sup>90,91</sup> These methods, however, do not provide strand-specific information, because the cDNA is subsequently fragmented and ligated to a second set of adapters. Recently, the multiple annealing and looping-based cycles (MALBAC) method, originally designed for single-cell genomic DNA amplification, was applied to RNA-Seq. 92

Because of a greater extent of amplification, molecular labels are particularly important for single-cell RNA-Seq to detect over-amplified products. In addition, barcoding is typically used to label cells, enabling simultaneous preparation of libraries from multiple cells. On this note, recent high throughput single-cell methods have used oligonucle-otide beads to deliver barcodes to cells and mRNAs at the same time (CytoSeq), or have separated cells with different barcodes in aqueous droplets (inDrop and Drop-seq). 94,95

State-of-the-art technologies are in development to characterize transcriptomes inside the cells, providing spatial information of RNA expression. One method, TIVA, is based on introduction of biotinylated tags to cells in tissue, followed by targeted activation of these tags by a laser in selected cells. The laser activates poly(U) tracts on the biotinylated tags, enabling them to bind mRNAs in the targeted cell. The bound mRNAs are then selected by streptavidin, cloned, and sequenced. Another method, fluorescent *in situ* RNA sequencing (FISSEQ), combines *in situ* amplification with sequencing by oligonucleotide ligation. 98

### SEQUENCING OF OTHER RNA SPECIES

### **Small RNAs**

Small noncoding RNAs below 30 nt, such as miR-NAs, piRNAs, and endosiRNAs, are processed from primary transcripts. miRNAs can be efficiently captured by direct ligation with adapters (see above).<sup>27</sup> Thanks to the 5' phosphate group and 3' OH group of miRNAs, no additional processing of RNA is necessary before the ligations. As discussed above, this method naturally preserves the strandedness of RNA but introduces substantial biases due to influence of sequence on ligation. Using degenerate random nucleotides at the ligation ends of adapters can effectively mitigate bias. 28,29 Alternatively, the 5' adapter ligation step, which appears to be more prone to bias than the 3' adapter ligation (personal observation), can be eliminated if the single-stranded cDNA is circularized by DNA ligase and amplified by PCR.99

Because of the short insert size of the cDNA library for small RNAs, it is necessary to use a specific method to separate them from contaminant DNAs, such as PCR products without any inserts. These methods include electrophoresis separation or adding the RT primer to block the 3' adapter from ligating with the 5' adapter.

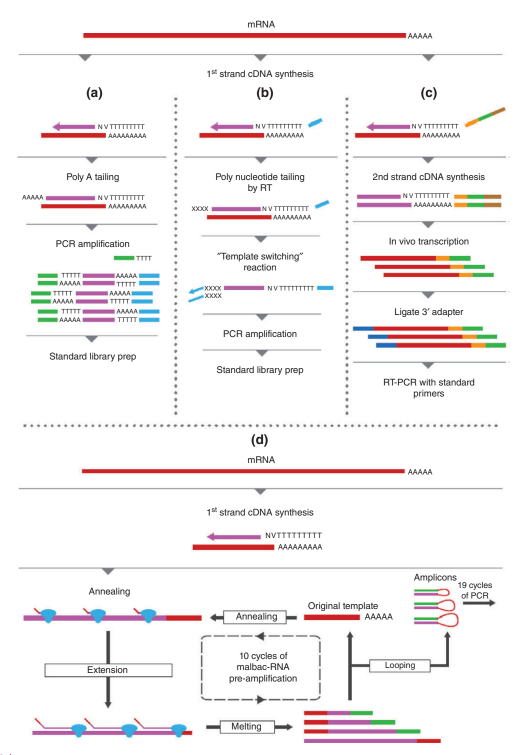
### Circular RNA

One recent surprising finding has been the discovery of circular RNAs,<sup>100</sup> which are generated by back-splicing.<sup>101</sup> Circular RNAs can be sequenced by digesting away linear RNAs using exonuclease R, followed by regular RNA-Seq methods involving fragmentation, RT and PCR.

### Specifically Generated RNA Fragments

A growing number of methods have been developed in the past few years to interrogate RNAs at different stages of biogenesis and metabolism or interactions with proteins or other RNAs, and RNA structural features. Some of these methods are summarized in Table 1. While these methods differ widely at the RNA-capturing step, ranging from RNase protection to immunoprecipitation to metabolic labeling, the cDNA library preparation methods are similar.

Most methods generate short RNA fragments that are often made into cDNAs using the small RNA sequencing approach (see above).



**FIGURE 4** | RNA-Seq of single cells. (a) Reverse transcription with oligo-dT primers and a universal primer sequence is followed by poly(A) tailing. After PCR amplifications, standard RNA-Seq libraries are prepared. (b) Reverse transcription incorporates a universal primer sequence. Template switching of reverse transcription is followed by annealing of the oligonucleotide with the sequence for a second PCR primer. (c) cDNA synthesis introduces the T7 promoter sequence at the 5′ end. After second strand cDNA synthesis, cRNA copies are generated by *in vitro* transcription. Finally, the second adapter is ligated to the 3′ end of the cRNA and libraries are constructed by PCR amplification. (d) Single-cell MALBAC RNA-Seq. Primers with seven random nucleotides at the 3′ end are annealed to cDNA and extended. Amplicons are looped to protect them from being further amplified. Ten cycles of quasilinear amplification are followed by exponential PCR.



**TABLE 1** Methods to Interrogate Different Aspects of RNA Life Cycle

Purpose	Methods	RNA Species Sequenced
Transcribing RNA polymerase	GRO-Seq <sup>102</sup>	Nascent RNA transcribed in vitro
	NET-Seq <sup>103–105</sup>	Nascent RNA associated with RNA polymerase in vivo
RNA synthesis, processing and degradation	4sU-Seq <sup>106–108</sup>	Newly synthesized RNA labeled with 4-thiouridine (4sU) in vivo
Translational status	Ribosome profiling (Ribo-seq) <sup>99</sup>	Ribosome-bound mRNA fragments protected from RNase digestion
RNA-protein interactions	RIP-Seq <sup>109–111</sup>	RNA in immunoprecipitated ribonucleoprotein complex
	HIT-CLIP/CLIP-seq, 112–114 iCLIP115	RNA fragments crosslinked to interacting proteins by UV (254 nm)
	PAR-CLIP <sup>116</sup>	RNA labeled with 4sU and crosslinked to interacting proteins by UV (365 nm)
RNA structure probing	PARS, <sup>117</sup> FragSeq <sup>118</sup>	RNA fragments generated by RNases digesting double- stranded regions (V1) or single-stranded regions (S1 or P1)
	DMS-Seq <sup>119</sup> ; SHAPE-S <sup>120</sup> icSHAPE <sup>121</sup>	Unstructured RNA regions labeled with reactive chemicals
RNA–RNA interactions	RAP-RNA <sup>122</sup>	RNA isolated by antisense probe

### CONCLUSION AND FUTURE DIRECTIONS

Since its inception about 8 years ago, RNA-Seq has become a widely used and indispensable approach for studying gene expression and interrogating aspects of RNA biogenesis and metabolism. As sequencing technologies continue to advance, many methods have been developed and new RNA-Seq methods are expected to emerge in the future. We believe several areas are particularly relevant to RNA analysis. First, current sequencing platforms have size limitations. Instruments that can handle long reads and offer high read output would be particularly beneficial for quantitative analysis of transcript

isoforms, including those generated by alternative initiation, AS, and alternative polyadenylation, and gene fusions. Second, current sequencing chemistry cannot handle homopolymers well. This is particularly relevant for sequencing the poly(A) tail region, which plays important roles in transcript metabolism. Third, the sensitivity of sequencing needs to be further improved so that amplification can be reduced or eliminated. This will also be important for single-cell analysis, where currently only a small fraction of genes can be examined. Finally, interrogation of spatial information of RNA expression in the cell, showing much promise in recent studies, is a new frontier for RNA-Seq.

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