Biogenesis, identification, and function of exonic circular RNAs

Iju Chen, Chia-Ying Chen and Trees-Juen Chuang

Circular RNAs (circRNAs) arise during post-transcriptional processes, in which a single-stranded RNA molecule forms a circle through covalent binding. Previously, circRNA products were often regarded to be splicing intermediates, by-products, or products of aberrant splicing. But recently, rapid advances in high-throughput RNA sequencing (RNA-seq) for global investigation of nonco-linear (NCL) RNAs, which comprised sequence segments that are topologically inconsistent with the reference genome, leads to renewed interest in this type of NCL RNA (i.e., circRNA), especially exonic circRNAs (ecircRNAs). Although the biogenesis and function of ecircRNAs are mostly unknown, some ecircRNAs are abundant, highly expressed, or evolutionarily conserved. Some ecircRNAs have been shown to affect microRNA regulation, and probably play roles in regulating parental gene transcription, cell proliferation, and RNA-binding proteins, indicating their functional potential for development as diagnostic tools. To date, thousands of ecircRNAs have been identified in multiple tissues/cell types from diverse species, through analyses of RNA-seq data. However, the detection of ecircRNA candidates involves several major challenges, including discrimination between ecircRNAs and other types of NCL RNAs (e.g., trans-spliced RNAs and genetic rearrangements); removal of sequencing errors, alignment errors, and in vitro artifacts; and the reconciliation of heterogeneous results arising from the use of different bioinformatics methods or sequencing data generated under different treatments. Such challenges may severely hamper the understanding of ecircRNAs. Herein, we review the biogenesis, identification, properties, and function of ecircRNAs, and discuss some unanswered questions regarding ecircRNAs. We also evaluate the accuracy (in terms of sensitivity and precision) of some well-known circRNA-detecting methods. © 2015 The Authors. WIREs RNA published by Wiley Periodicals, Inc.

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INTRODUCTION
Following the discovery of the first circular RNA (circRNA) molecules,1 several types of RNA circles have been detected in various organisms. Unlike plant viroids and the hepatitis delta virus,1,2 which have circular single-stranded RNA (ssRNA) genomes, several transcribed RNA molecules, including tRNAs, rRNAs, and mRNAs, can also be circularized via ribozymal activity (Group I3 and Group II4 introns), archaeal splicing,5,6, or spliceosomal machinery.7–9 In the past, observed circRNAs (especially in animals) were usually thought to be by-products of pre-mRNA processing, and were therefore interpreted to be results of missplicing.7–9 Recently, advances in high-throughput RNA sequencing (RNA-seq) have created unprecedented opportunities to globally investigate transcriptomes, revealing the existence of a large amount of previously
unidentified circRNAs, especially of exonic circRNAs (ecircRNAs).\textsuperscript{10} Genome-wide analysis of RNA-seq data revealed that ecircRNAs are abundant in mammalian transcriptomes, and some of them are evolutionarily conserved in terms of sequence and expression,\textsuperscript{10–12} suggesting that they possess cellular function. The most prominent examples of functional ecircRNAs are human \textit{CDR1as/ciRS-7} and circRNA of mouse \textit{Sry}, which were experimentally validated to function as miRNA sponges, and are thereby involved in gene expression regulation.\textsuperscript{13,14} Moreover, some ecircRNAs were originated from genes related to splicing factors,\textsuperscript{15} DNA methyltransferases,\textsuperscript{15} and important diseases such as dystrophy\textsuperscript{16} and cancers.\textsuperscript{7,9} Although the relationship between ecircRNAs and their linear counterparts is mostly unknown, an understanding of such an association with disease-forming processes will help considerably in developing efficient diagnostic methods or even therapies. Of particular note, ecircRNAs were shown to be more stable than their linear counterparts in plasma\textsuperscript{17} and saliva,\textsuperscript{18} suggesting their potential as diagnostic biomarkers.

Generally, an ecircRNA event can be detected by aligning expressed sequences (e.g., RNA-seq reads) against the reference genome. After that, non-co-linear (NCL) junctions are determined on the basis of the presence of two connected exons in which the exon order is topologically inconsistent with the reference genome. Although varied bioinformatics methods based on RNA-seq data have been developed and used to identify thousands of ecircRNA candidates in diverse species (Table 1), there remain several challenges. For example, an observed NCL junction site may also be formed by other types of NCL event (e.g., \textit{trans}-splicing events and genetic rearrangements) or various types of false positives (e.g., sequencing errors, alignment errors, and \textit{in vitro} artifacts). In addition, the identification results may drastically differ between different circRNA-detecting methods and data derived from different RNA treatments, resulting in biased interpretations for ecircRNA analysis. To date, there is no competent method that can simultaneously account for the abovementioned issues during the identification of NCL events (including circRNAs). Moreover, there remains a need to evaluate the sensitivity and precision of currently available methods for circRNA identification.

In this review, we focus on the discussion of the biogenesis, identification, properties, and function of ecircRNAs. We also describe our generation of

### Table 1: Recently Published Studies for Detecting circRNAs on the Basis of RNA Sequencing Data

| Study               | Exonic/intronic circRNA | Treatment of RNA Library | Number of Detected circRNA Events | Method for circRNA Identification |
|---------------------|-------------------------|--------------------------|-----------------------------------|----------------------------------|
| Salzman et al. (2012)\textsuperscript{10} | Exonic                   | rRNA\textsuperscript{−} | >880 human genes and >1000 mouse genes contain circRNAs | In-house pipeline |
| Salzman et al. (2013)\textsuperscript{19} | Exonic                   | rRNA\textsuperscript{−}\&polyA\textsuperscript{−} | 46,866 events in 8466 human genes | In-house pipeline |
| Zhang et al. (2013)\textsuperscript{20}   | Intronic                | rRNA\textsuperscript{−}\&polyA\textsuperscript{−}; rRNA\textsuperscript{−}\&RNase R\textsuperscript{+} | 103 events (human) | The in-house pipeline |
| Zhang et al. (2014)\textsuperscript{15}   | Exonic                  | rRNA\textsuperscript{−}\&polyA\textsuperscript{−}; rRNA\textsuperscript{−}\&RNase R\textsuperscript{+} | 1662 events (human) | CIRCexplorer |
| Jeck et al. (2013)\textsuperscript{11}   | Exonic                  | rRNA\textsuperscript{−}\&RNase R\textsuperscript{+} | 7771 events (human); 646 events (mouse) | MapSplice |
| Memczak et al. (2013)\textsuperscript{13} | Exonic                  | rRNA\textsuperscript{−} | 1903 events (human); 1111 events (nematode) | find_circ |
| Ashwal et al. (2014)\textsuperscript{21} | Exonic                  | rRNA\textsuperscript{−}\&RNase R\textsuperscript{+} | 3117 events (fruit fly) | In-house pipeline |
| Hoffmann et al. (2014)\textsuperscript{22} | Exonic                  | rRNA\textsuperscript{−} | 1712 events (human) | segemehl |
| Guo et al. (2014)\textsuperscript{23}    | Exonic                  | rRNA\textsuperscript{−} | 7112 events (human); 635 events (mouse) | In-house pipeline |
| Westholm et al. (2014)\textsuperscript{24} | Exonic                  | rRNA\textsuperscript{−} | 2513 (fruit fly) | In-house pipeline |
| Bachmayr-Heyda et al. (2015)\textsuperscript{25} | Exonic                  | rRNA\textsuperscript{−} | 1812 (human) | find_circ |
| Gao et al. (2015)\textsuperscript{26}     | Both                    | rRNA\textsuperscript{−} | 3000–10,000 (human) | CIRI |
artificial paired-end RNA-seq reads from a mix of simulated intragenic NCL transcripts and well-annotated co-linear transcripts to evaluate the sensitivity and precision of five well-known circRNA-detecting tools: TopHat-Fusion,\textsuperscript{27} MapSplice,\textsuperscript{28} seqemehl,\textsuperscript{22} find_circ,\textsuperscript{13} and CIRI.\textsuperscript{26} Some unanswered questions regarding ecircRNA biogenesis, identification, and function are also discussed.

**BIOGENESIS OF EXONIC CIRCORNAS**

**Models of ecircRNA Formation**

Eukaryotic exonic regions of pre-mRNAs are typically disrupted by intron(s), and spliceosomes are responsible for the removal of introns from pre-mRNAs. A generally accepted model depicts a two-step spliceosome action:\textsuperscript{29} (1) the branch point (BP, usually an adenosine) attacks the 5′ splice site (5′SS) via its 2′-hydroxyl group, forming a 2′-5′ phosphodiester bond and a free 3′-hydroxyl end on the 5′ exon; and (2) the newly generated 3′-hydroxyl attacks the 3′ splice site (3′SS), and a 3′-5′ phosphodiester bond is formed to ligate the two exons; meanwhile, the intron lariat with a 2′-5′ linkage is excised. Strict control of the choice of splice site pairs is important, as it ensures the accuracy of mRNA products and consequent processes. However, splicing machinery also shows certain degrees of flexibility (sometimes described as aberrant) in splice site choice. Variance in splice site pairing (or alternative splicing) often creates varied transcript isoforms. As many ecircRNAs contain exons from coding genes and are connected by canonical splice sites, it is generally believed that ecircRNAs are produced through spliceosomal splicing mechanisms. Some evidence has been proposed to support this scenario. One line of direct evidence supporting the essential role of the spliceosomal machinery in ecircRNA biogenesis comes from the use of splice inhibitor isoginkgetin.\textsuperscript{30} Following isoginkgetin treatment, both linear and circular isoforms were significantly reduced in nascent RNA pools. Mutagenesis analyses show that both 5′ and 3′ splice signals from the circular junction are essential for exon circularization.\textsuperscript{21,30} These results supported the hypothesis that spliceosome-mediated pre-mRNA splicing may involve backsplicing (or reverse splicing), which connects a downstream splice donor site (5′ splice site) to an upstream acceptor splice site (3′ splice site) and forms an ecircRNA. Comparison of backsplicing and canonical (or linear) splicing further indicated that although canonical splicing factors can control both processes, the splicing regulatory rules for circRNA biogenesis are different from those for linear splicing.\textsuperscript{31} In addition, it was proposed that linear splicing and circularization may compete for limited splicing factors—introducing flanking exons with strong 5′ and 3′ splice sites dramatically decreases circularization efficiency.\textsuperscript{21} In addition, the minigene system, which is frequently used to investigate the splicing mechanism, is also used to investigate ecircRNA biogenesis. A recent study showed that, in addition to canonical splice signals, important signal sequences in the spliceosomal machinery (such as poly-pyrimidine tracts) also influence circularization; however, the involvement of the branch point is less conclusive.\textsuperscript{30} Other studies demonstrated that changes in encircled exonic sequences can abolish circRNA formation,\textsuperscript{32} but in some cases it does not affect circularization.\textsuperscript{30} Several models that were proposed to explain the possible formation of ecircRNAs are discussed below (see also Figure 1).

**Lariat-driven Circularization**

In an exon skipping (cassette-on) event, the spliced intron lariat also contains the skipped exon(s) (Figure 1(a)). If further splicing occurs within the lariat before the unraveling of the lariat by debranching enzymes, a stable RNA circle enclosing the skipped exons can be generated.\textsuperscript{11} Meanwhile, a linear transcript excluding the skipped exon(s) is also produced. Exon skipping was suggested to be the cause of circRNA formation in some early cases since the linear counterparts of such skipping events were detected.\textsuperscript{33,34} Genome-wide analysis of RNA-seq data from a human fibroblast cell line revealed that, for 45% of 7771 predicted circRNAs, the corresponding linear isoforms also exhibited exon skipping events,\textsuperscript{11} suggesting that RNA circularization was correlated with exon skipping. However, such a trend was not observed in a separate study on different biosamples.\textsuperscript{10}

**Intron Pairing-driven Circularization**

In this model, the formation of ecircRNAs is independent of exon skipping. It differs from the lariat-driven circularization model by the choice of splice site pairs and the lack of knowledge about the corresponding linear product(s). It was suggested that intronic motifs might border the circularized exon(s) and thereby join the circularized exon(s)\textsuperscript{11} (Figure 1(b)). Distinguishing between lariat- and intron pairing-driven ecircRNAs is difficult, because the corresponding products/intermediates are likely to be short-lived, as a result of degradation through nonsense-mediated decay or by debranching enzymes.

**Resplicing-driven Circularization**

In the presence of proper cis- and trans-splicing elements, resplicing may take place on spliced mRNAs.
FIGURE 1 | Possible models of ecircRNA biogenesis. (a) Lariat-driven circularization, (b) Intron-pairing-driven circularization, and (c) Resplicing-driven circularization. B, branch point.

Criteria for Exon Circularization

The Involvement of Reverse Complement Sequences

It has been shown that base pairing between the reverse complementary sequences (RCSs) in flanking introns can bring the downstream 5’SS into the proximity of the upstream 3’SS (Figure 2(a)), leading to circularization of the mouse Sry gene.\(^8\),\(^36\) Several transcriptome-wide analyses also indicated a significant correlation between the presence of flanking intronic RCSs (especially inverted Alu elements in primates; Figure 2(a)) and exon circularization.\(^11\),\(^15\),\(^37\) Extensive mutagenesis of expression plasmids revealed that short (30–40 nt) inverted repeats (e.g., Alu elements) are sufficient for ecircRNA generation.\(^32\) However, it was observed that not all intronic repeats could support exon circularization; on the contrary, enhancing the stability of base-pairing sometimes might impede circRNA formation.\(^32\) In addition, if multiple copies of RCSs are present in a single gene, the competition for base pairing among RCSs may affect circularization efficiency, and even result in alternative circularization, bringing more diversity of circular transcripts from a single gene.\(^15\) It was demonstrated that circRNAs can be predicted by scoring the presence of RCSs in the bracketing introns.\(^37\) Although exon circularization is highly correlated with intronic RCSs, several studies using the minigene system showed that human ecircRNAs are not always bracketed by Alu or RCSs-containing introns, further indicating that RCSs can enhance,\(^15\),\(^30\)\(–32\) but are not essential for,\(^30\),\(^31\) ecircRNA production. The bracketing introns of ecircRNAs are also highly enriched for RCSs in animals that are not rich in repeats, such as Caenorhabditis elegans\(^37\) and Drosophila\(^24\),\(^31\); however, it is formally
**Possible that ecircRNA biogenesis may be regulated by different cis elements in different species.**

**Regulatory Factors for Circularization**
Although ecircRNA biogenesis can be viewed as a mode of alternative splicing, it remains unknown whether factors involved in alternative splicing regulation are also involved in ecircRNA biogenesis. A recent study appeared to partially answer this question by demonstrating that a considerable number of ecircRNAs are dynamically regulated by Quaking (QKI), an alternative splicing factor, during the human epithelial-mesenchymal transition.38 The addition of QKI binding motifs to flanking introns can significantly induce circRNA formation,38 suggesting that QKI is an important regulator of circularization. Another regulator of circRNA biogenesis is muscleblind (MBL/MBNL1), a splicing factor that was found to be circularized in flies and humans.21 This circRNA contains multiple MBL-binding motifs in its flanking introns, which are specifically bound by MBL. Downregulation of MBL can result in a remarkable decrease in circularization.21 Both QKI and MBL promote ecircRNA generation by bringing the 5′ SS closer to the upstream 3′ SS (Figure 2(b)). On the other hand, the double-strand RNA-editing enzyme – adenosine deaminase acting on RNA (ADAR) proteins, which tend to bind and mediate A-to-I editing on inverted Alu repeats, were also demonstrated to regulate circRNA biogenesis.37 Disruption of ADAR expression could result in a significant increase in circRNA expression in C. elegans and human,12,37 suggesting that ADAR proteins might play an antagonistic role in circRNA production (Figure 2(c)).

**IDENTIFICATION OF EXONIC CIRC RNAS**

**Strategies to Identify ecircRNAs**
Many RNA-seq-based bioinformatics tools have been developed to identify ecircRNA candidates. Table 1 summarizes some recently published studies on the detection of ecircRNAs. Basically, ecircRNAs are detected by comparing the reference genomes with RNA-seq reads, and then extracting matches comprised of sequence segments topologically inconsistent with the corresponding DNA sequences in the reference genome. According to the dependency on genome annotation (i.e., annotated exon–intron boundaries), these tools can be classified into two categories: pseudo-reference- and fragment-based strategies (Figure 3 and Table 1). For the pseudo-reference-based strategy, genome annotation is required. All possible combinations of pseudo references are constructed; each of them is comprised of two well-annotated exons in which the exon order is topologically inconsistent with the reference genome (Figure 3(a)). A pseudo reference is regarded as a circRNA candidate if it has at least one read that maps to its NCL junction site (Figure 3(a)). On the other hand, the fragment-based strategy detects circRNAs without the help of genome annotation. RNA-seq reads (each paired-end read is viewed as two ‘single’ reads) are split into two or more segments, and each segment is mapped to the reference genome; segmented reads mapped in an NCL manner are retained (Figure 3(b)). There are two major limitations for pseudo-reference-based methods: first, they cannot identify circRNAs with unannotated...
FIGURE 3 | Two RNA-seq-based strategies for detecting NCL junctions of ecircRNA candidates: (A) pseudo-reference-based and (B) fragment-based strategies. The former identifies NCL junction sites at annotated exon junctions; whereas the latter does not. NCL, non-circular.

exon junctions; and second, they are not suitable for detection of circRNAs in the genomes that are incomplete or poorly annotated. On the other hand, the fragment-based methods can be used to identify NCL junctions at a single nucleotide resolution in the absence of any existing genome annotation. However, as segmented reads are smaller than full-length reads, such an approach is more likely to yield alignment errors (or ambiguity) than the pseudo-reference-based strategy while performing read-to-genome alignment. In addition, NCL junctions that do not match annotated exon boundaries tend to be unreliable and are more likely to originate from missplicing.39–42

Certain additional criteria are often applied to improve the accuracy of ecircRNA identification. For example, if paired-end reads are used to identify ecircRNAs, both ends of each matched read should (i) be mapped to the circle predicted by the circular junction and (ii) be in the correct orientation (Figure 4(a)). From the mapping patterns of paired-end reads, various scenarios for circles, such as circles containing a single exon, partial fragment(s) of an annotated exon, or an intron-containing fragment, can be depicted (Figure 4(b)). In addition, fragment-based strategies generally consider only NCL junctions that are flanked by the GT-AG canonical splice sites for improving accuracy.

Moreover, the observations that circRNAs are non-polyadenylated10,19,23 or RNase R-resistant11,13,43 have been exploited by many studies to increase the accuracy of ecircRNA identification (Table 1). Circular-junction candidates are often detected from an rRNA-depleted total RNA library, and filtered by comparison to candidates detected in other library sets treated with either RNase R6,11,20 or poly(dT).20,24 However, although such approaches detect abundant circles, the following matters need to be borne in mind. First, not all backsplicing events show enrichment in RNase R-treated RNA libraries. For example, CDR1as/ciRS-7 was not enriched after RNase R treatment.11 Second, circRNAs and trans-splicing events sometimes share the same NCL junctions,41 and such junctions are therefore present in both poly(A)- and non-poly(A)-selected RNA-seq data. Third, not all mRNAs lacking poly(A) are circular; for example, certain replication-dependent histone genes are co-linear transcripts without poly(A) tails.44 Fourth, the amount of circRNA events are often amplified in the treated RNA-seq data as compared to the data from untreated samples, raising the concern that a considerable number of detected events may represent rarely but pervasively occurring ‘background’ NCL junctions derived from splicing errors.11 Finally, such treated data are sensitive to endonuclease contamination, and may also be less effective for identifying longer exons.11,45,46

Difficulties of ecircRNA Identification

Identification of ecircRNAs often suffers as a result of three major challenges: (1) discrimination between
circRNAs and other types of NCL events, such as trans-splicing and genetic rearrangements; (2) removal of false positives arising from sequencing errors, alignment errors, and in vitro artifacts; and (3) biased identification of circRNAs from different bioinformatics methods or the use of sequencing data from different treatments. In fact, these difficulties are common in the detection of all types of NCL RNAs.  

**Removal of False Positives**

As stated above, circRNAs are one class of NCL RNAs. Detecting all types of NCL event often suffers from false positives arising from sequencing errors, alignment errors, and in vitro artifacts. These false positives can severely affect the accuracy of detecting NCL RNAs. In general, false positives caused by sequencing errors can be reduced by increasing the number of RNA-seq reads that support the NCL junctions, or by eliminating skew mapping between reads and the corresponding NCL junctions. However, as most circRNAs are expressed at a relatively lower level compared with co-linear mRNAs, such approaches may sacrifice a considerable number of true positives unless the sequencing depth is very deep. Furthermore, as paralogous genes or repetitive sequences are prevalent in genomes, ambiguous alignments during short-read mapping are often misinterpreted as NCL events. In particular, sequencing errors within repetitive sequences can increase the chances of mapping errors, and result in misidentified backsplcing junctions. A recent study suggested that comparison of different alignment results can effectively eliminate ambiguous alignments. Nevertheless, it is still very difficult to determine whether an observed NCL junction arises from ambiguous alignments in incomplete or draft genomes. For circRNA detection, a previous study eliminated potential alignment errors by controlling for the alignment quality of both ends of RNA-seq reads that were mapped inside a circle candidate. However, it remains a major challenge to effectively remove alignment errors without losing sensitivity.

Finally, spurious NCL events may also be generated from artificial RNA-seq reads that are produced during cDNA library construction. As RNA-seq data are generally derived from reverse transcriptase (RT)-based sequencing approaches, RT artifacts, such as template switching, often impede the accurate identification of NCL events. Reverse transcriptase may switch templates in the process of reverse transcription, either to a different RNA molecule or to a different location on the same template. Several studies have demonstrated that the majority of NCL events extracted from mRNAs were generated from experimental artifacts. Unfortunately, it is difficult to distinguish such artifacts from genuine NCL events by simple experimental validations, not mention to the NCL RNA candidates merely identified by bioinformatics strategies without any experimental validation. Recently, some NCL events that previously passed RT-PCR validations were subsequently confirmed by more careful validations to be originated from
in vitro artifacts.\textsuperscript{41} Previous studies have indicated that increasing the primer annealing temperature during reverse transcription may reduce the emergence of template switching events.\textsuperscript{53,54} However, such experiments were shown to be insufficient to eliminate template switching-derived NCL events.\textsuperscript{40,51} It was also demonstrated that such RT artifacts cannot be easily removed by controlling for canonical splice signals encompassing the NCL junctions or the depth of RNA-seq reads supporting the NCL junctions.\textsuperscript{40,41,52} Several studies demonstrated that RTase-dependent RNA products were likely to be RT artifacts, suggesting that comparisons of different RTases products could effectively detect such artifacts.\textsuperscript{40,41,51} Alternatively, some non-RTase-based experiments, such as Northern blot and RNase protection assay,\textsuperscript{55} can be applied to the detection of RT artifacts, although these validations are more expensive and time consuming than RTase-based ones. To date, there is only one systematic approach that can detect NCL RNAs while controlling for experimental artifacts.\textsuperscript{52} Unfortunately, this approach is based on \textit{Drosophila} hybrid mRNAs (\textit{Drosophila melanogaster} females vs. \textit{Drosophila sechellia} males) and a mixed mRNA-negative control sample,\textsuperscript{52} and thus cannot be applied to human studies.

\textbf{Biased Identification of circRNAs}

There are many discrepancies among circRNA candidates identified by different methods,\textsuperscript{27} with the major contributing factor being that different methods used different detection rules to identify circRNAs.\textsuperscript{56} Such discrepancies between results also imply that a considerable proportion of detected circRNA candidates are merely false positives. In addition, using RNA-seq data derived using different RNA-library treatments to detect circRNA candidates may also yield different results. To examine this issue, we used five well-known circRNA-detecting methods, TopHat-Fusion,\textsuperscript{27} MapSplice,\textsuperscript{28} segemehl,\textsuperscript{22} find\_circ,\textsuperscript{13} and CIRI,\textsuperscript{26} to individually detect circRNA candidates in HeLa cells with three different RNA-library treatments (Table 2): rRNA depletion (rRNA\textsuperscript{−}), rRNA-depleted RNAs with RNase R treatment (rRNA\textsuperscript{−}\&RNase R\textsuperscript{+}), and rRNA-depleted RNAs with poly(A) depletion (rRNA\textsuperscript{−}\&polyA\textsuperscript{−}). Of note, as TopHat-Fusion, MapSplice, and segemehl can also detect intergenic NCL events, we only considered the intragenic NCL events detected by these three methods. We showed that a considerable proportion (23–85\%) of the detected circRNA candidates is dependent on the individual RNA-library treatment (Figure 5(a)). Such proportions vary among the methods used (Figure 5(a)), which also reflect the discrepancies among different identification results. This result thus indicates that genome-wide analysis of circRNAs may be biased by both the method and RNA-library treatment. Moreover, we find that as much as 31–76\% of the intragenic NCL events detected in rRNA\textsuperscript{−} data are absent from both poly(A)-depleted data and RNase R-selected data (Figure 5(a)). As circRNAs tend to be non-polyadenylated\textsuperscript{10,19,23} or RNase R-resistant,\textsuperscript{11,13,43} such intragenic NCL events that are dependent on RNA-depleted data may not arise from backsplicing, suggesting that these circRNA candidates should be further curated.

\textbf{Evaluation of Sensitivity and Precision of circRNA-Detecting Methods}

To evaluate the sensitivity (\(S_n\)) and precision (\(S_p\)) of different circRNA-detecting methods, we utilized Mason\textsuperscript{17} to generate paired-end reads (with read length 2 \(\times\) 100 nt) from 100 simulated intragenic NCL transcripts with different expression levels (5- to 100-fold), and then mixed these simulated data with the same background dataset generated from the GENCODE-annotated (version 19) co-linear transcripts. The simulated NCL transcripts must not be derived from pseudogenes or mitochondrial or ribosomal genes,\textsuperscript{58} and their junction sites were randomly generated and located at the boundaries of annotated exons. The above-mentioned circRNA-detecting methods (i.e., TopHat-Fusion, MapSplice, segemehl, find\_circ, and CIRI) were then applied to the simulated datasets. Our results revealed that the \(S_n\) and \(S_p\) values were both positively correlated with the expression levels of circRNAs (Figure 5(b)). When examining the tested dataset at all simulated expression levels, TopHat-Fusion exhibited the highest \(S_n\) values but the

| RNA-library Treatment | Sequencing Platform | Read Length | NCBI SRA ID (Read Number) |
|-----------------------|---------------------|-------------|---------------------------|
| rRNA\textsuperscript{−} | Illumina Hisq 2000 | 101 bp      | SRR1637089 (44,933,450) SRR1637090 (35,685,310) |
| rRNA\textsuperscript{−}\&RNase R\textsuperscript{+} | Illumina Hisq 2000 | 101 bp      | SRR1636985 (13,309,745) SRR1636986 (23,505,713) |
| rRNA\textsuperscript{−}\&polyA\textsuperscript{−} | Illumina GA II | 76 bp       | SRR317048 (70,788,979) |
FIGURE 5 | Comparison of identified ecircRNAs based on different bioinformatics methods or the use of sequencing data from different treatments. (a) Venn diagram of identified circRNAs based on HeLa cell transcriptome data with different RNA-library treatments for each individual algorithm. The percentage of ecircRNA events identified from each RNA-library treatment is showed in parentheses. (b) Evaluation of sensitivity ($S_n$) and precision ($S_p$) of five ecircRNA-detecting algorithms, based on simulated datasets of different expression levels of NCL transcripts. $S_n$ and $S_p$, both of which range from 0 to 1, are defined as TP/(TP+FN) and TP/(TP+FP), respectively. TP (true positive), FP (false positive), and FN (false negative) represent the number of correctly identified events, the number of incorrectly identified events, and the number of missing events, respectively.
lowest \( S_p \) values (all \( S_p < 0.1 \)), whereas the opposite was observed for MapSplice (Figure 5(b)). It was notable that although MapSplice achieved 100% precision (all \( S_p = 1 \)) under all simulated conditions, it had very poor sensitivity (all \( S_p < 0.4 \)) (Figure 5(b)). This reveals that certain methods achieve better precision by sacrificing sensitivity, highlighting the difficulty in reaching a balance between sensitivity and precision. Overall, segemehl, find_circ, and CIRI exhibited similar levels of sensitivity, but find_circ and CIRI demonstrated relatively lower precision (all \( S_p < 0.3 \)) than segemehl (all \( S_p > 0.6 \)) (Figure 5(b)). Therefore, here segemehl seemed to achieve a better balance between sensitivity and precision than the other methods examined. Of note, here all tools for evaluation were used with default parameters. In fact, different stringency levels of parameter settings (e.g., the number of RNA-seq reads supporting the NCL junctions, the alignment quality of both ends of mapped RNA-seq reads, the control of canonical splice signals encompassing the NCL junctions, etc.) may significantly affect the number of identified candidate circles for the same tool, and thereby affect the accuracy. Generally, circRNA-detecting tools with low-stringency parameters could achieve better sensitivity but worse precision than those with high-stringency ones.

**GLOBAL PROPERTIES OF EXONIC CIRCULAR RNAs**

**Flanking Introns of ecircRNAs**

In addition to the aforementioned excess of RCSs in ecircRNA-flanking introns, various independent studies reached the following conclusion: ecircRNAs tend to have longer flanking introns than expected (comparing to the average or control sets) in diverse species (e.g., humans, flies, and nematodes).\(^{10,11,15,24,37}\) In human, the ecircRNA-flanking introns are three- to fivefold longer than randomly selected introns.\(^{11,15}\) In *Drosophila*, upstream and downstream flanking introns of ecircRNAs have median lengths of 4662 and 2962 nt, respectively, both of which are much longer than the median length of all introns (94 nt).\(^{24}\) In addition, *C. elegans* ecircRNAs were observed to have 10-fold longer flanking introns than the median length of all the introns.\(^{37}\) Although longer flanking introns seemed to promote ecircRNA formation, statistical analysis indicated that long flanking introns were not necessary for ecircRNA formation in humans.\(^{19}\) A later study revealed that a longer flanking intron itself does not cause ecircRNA formation; instead, the longer the intron, the greater the possibility that it contains more *cis* elements (e.g., inverted *Alu* elements) that promote ecircRNA formation.\(^{15}\) However, no specific motifs or structures have been found in flanking intron pairs so far.\(^{24}\) It is known that different species exhibit remarkable variations in intron length. Whether such a fundamental difference may have influenced ecircRNA biogenesis awaits further elucidation.

**Sequence Context of ecircRNAs**

Recent transcriptome-wide analyses have revealed several common features in animal ecircRNAs. First, ecircRNAs may consist of a single exon or multiple exons (see also Figure 4(b)). Many of ecircRNAs encircle the second exon\(^{10}\) or the exon(s) near the 5′ end\(^{24}\) of the corresponding co-linear counterpart. Typically, one gene contains one circular form, but some genes can form multiple circular products. Most human ecircRNAs contain less than 5 exons with a median length of 547 nt.\(^{23}\) Only a few ecircRNAs are smaller than 80 nt in length.\(^{26}\) Long exons tend to be enclosed within the circles.\(^{11}\) For the case of circRNAs with a single exon, the encircled exons are longer than overall expressed exons.\(^{11,15}\) Second, GT-AG canonical splice sites are usually required for circRNA formation, although cryptic splice sites are sometimes used instead.\(^{30}\) Splice sites used for co-linear *cis*-splicing\(^{10,11,24}\) or NCL *trans*-splicing\(^{41}\) may also be used to form ecircRNAs. Nevertheless, there is no special global pattern associated with splice sites for circRNA biogenesis.\(^{30}\) Third, introns between the encircled exons are usually excised, but are retained in some rare cases (Figure 4(b)).\(^{11,23,24}\) Noncoding RNAs, intergenic regions, or antisense regions are possibly (if not seldom) encompassed in ecircRNAs.\(^{23,24,26}\) Fourth, with the exception of the splice site sequences, RNA motifs are generally not present within the circles shared by most ecircRNAs. Only a handful of ecircRNAs were observed to contain MBL motifs\(^{21}\) and miRNA binding sites.\(^{13,14}\)

**Conservation of ecircRNAs among Species**

Several studies have shown that some ecircRNAs are evolutionarily conserved between three *Drosophila* species\(^{23,24}\) or between humans and mice,\(^{11,13,23,37}\) implying circular forms are not by-products of splicing or randomly misspliced products. The conservation of splicing regulatory elements in host genes may be responsible for the conservation of circRNA generation between species. A recent study revealed that the ecircRNAs expressed in both human and mouse have a higher probability of forming circles by base-pairing of RCSs in the flanking introns as compared with those
expressed only in one species or exons with bracketing intron length-matched controls, suggesting that the human–mouse orthologous ecircRNAs are also conserved in terms of their exon circularization between these two species. However, exons within the human–mouse orthologous circles do not exhibit greater sequence conservation than their neighboring linear exons. \( ^{23} \) The correlation between retention of ecircRNA orthologues across evolutionarily distant species and biological significance demands further investigation.

**Abundance and Tissue-specific Accumulation**

A large amount of ecircRNAs have been identified in various cell types and eukaryotes (Table 1). A prominent website, circBase,\(^ {59} \) continues to collect identified circRNAs. In human, as high as \( \sim 100,000 \) ecircRNAs have been identified (Table 1), although the number of human ecircRNAs should be estimated more conservatively because of the possibility of false positives arising from high-throughput sequencing or identification processes (as described above). It was suggested that circRNAs comprise 1 to \( \sim 10\% \) of all transcripts in human cells.\(^ {10,19,23} \) No specific pattern of association between the circular forms and their corresponding linear transcripts has been observed. Most ecircRNAs are expressed at a very low level (0.1–1% of the expression levels of their co-linear counterparts), but a few cases were more abundant than their co-linear isoforms.\(^ {11} \) When it comes to tissue-specificity, most circRNAs were detected in only a few tissues/cell types.\(^ {13,19,24,26,30} \) A study used 15 human cell types to show that widely expressed circRNAs exhibited significantly higher expression than narrowly expressed ones.\(^ {26} \) Interestingly, in flies, ecircRNAs tended to arise from neural-related genes and had a higher expression level in neural tissues.\(^ {24} \) Similarly, ecircRNAs were reported to be enriched in mouse brain.\(^ {60} \)

**Subcellular Localization**

Several studies have shown that ecircRNAs are enriched in cytoplasmic samples.\(^ {10,11,13,23} \) As ecircRNAs are generated by the spliceosomal machinery in the nucleus and can be found in chromatin-bound RNA pools, they are likely to be transported by the nuclear export system, or escape from nuclei during cell division. Cytosolic localization may also support the post-transcriptional function of ecircRNAs. The most representative examples are \( CDR1as/ciRS-7 \) and circRNA Sry, which are predominantly localized in the cytoplasm and function as miRNA sponges when the specific miRNA (miR-7 for \( CDR1as/ciRS-7; \) and miR138 for circRNA Sry) is present.\(^ {13,14} \) However, a very recent study showed that some ecircRNAs with intronic segments retained between exons were predominantly located in the nucleus, where they \( cis \)-regulated their parent genes through specific RNA–RNA interactions.\(^ {61} \) These observations indicate that different ecircRNAs may differ in their preferred subcellular localizations, suggesting they may also possess varied functions.

**Translation Potential**

As most ecircRNAs carry open reading frames, one may speculate that they may be translated into peptides. It was shown that peptides can be translated from ecircRNAs \textit{in vitro} or \textit{in vivo},\(^ {26} \) as initiated from viral internal ribosome entry sites (IRESs)\(^ {62} \) or from prokaryotic ribosome-binding sites.\(^ {63} \) Translation of ecircRNAs produced from backsplicing in human cells transfected with vectors has recently been demonstrated.\(^ {31} \) Nevertheless, there is no evidence that spliceosome-generated ecircRNAs can serve as mRNAs. Analyses based on mass spectrometry data,\(^ {19,60} \) ribosome profiling,\(^ {23,60} \) and polysome profiling\(^ {8,11,60} \) have indicated that ecircRNAs tend to be untranslatable.

**FUNCTIONS OF EXONIC CIRCULAR RNAs**

**miRNA Sponge**

The ecircRNAs from human/mouse \( CDR1as/ciRS-7 \) and mouse Sry have been experimentally validated to be highly associated with the miRNA effector protein Argonaute in the presence of miR-7 and miR-138, respectively.\(^ {13,14} \) \( CDR1as/ciRS-7 \) contains 74 miR-7 binding sites, while circRNA Sry contains 16 miR-138 binding sites. The miRNA binding does not destabilize these two ecircRNAs; instead it competes with the binding between the miRNA and its target coding genes, and thereby reduces the effect of miRNA-mediated posttranscriptional repression. It has been conclusively demonstrated that over-expressing circRNAs of \( CDR1as/ciRS-7 \) or Sry increases the expression of miRNA target reporter constructs, while knockdown of these ecircRNAs has the opposite effect.\(^ {14} \) Downregulation of miR-7 targets was also observed in \( CDR1as/ciRS-7 \) knockdown human cells.\(^ {13} \) Therefore, these two ecircRNAs are believed to serve as miRNA sponges\(^ {13,14} \) or miRNA reservoirs\(^ {64} \) to attenuate miRNA-mediated responses.
In other words, with the same specific miRNA binding sites, ecircRNAs may play a regulatory role for competing endogenous RNA activities, which act as miRNA decoys to regulate the miRNA effects on their coding RNA targets by competing for miRNA binding. Expression of the corresponding coding RNA targets could be elevated by increasing the expression of such competing endogenous RNAs (or ‘ceRNAs’). An in vivo experiment in zebrafish further demonstrated that injection of human or mouse CDR1as/ciRS-7 could lead to reduced midbrain sizes, similar to the phenotype of miR-7 knockdown.13 In addition to miR-7 miRNA sites, CDR1as/ciRS-7 carries one miR-671 site, which triggers its own linearization and destruction.65 Moreover, ecircRNAs from the human C2H2 zinc finger gene family were also predicted to function as miRNA sponges.23 A bioinformatics study showed that the miRNA sites in circRNAs are depleted of polymorphisms, suggesting the important role of circRNAs in regulating miRNA activities.66 Although these results indicated that some circRNAs indeed function as miRNA sponges, several studies suggested that most circRNAs do not act as miRNA-sponges, as a large majority do not have more miRNA binding sites than co-linear mRNAs.23,60

Regulatory Role of ecircRNAs during Development and Cell Proliferation

It was observed that ecircRNAs tended to be tissue specific and enriched in brain.12,24,60 Although the reason may be that most the parent genes of ecircRNAs were also enriched in brain, the relative contribution of ecircRNA to the total transcriptional output of the same gene was remarkably higher in brain than in other tested tissues.60 Compared with the corresponding co-linear isoforms, ecircRNAs increase during aging of the central nervous system24 and decrease during cell proliferation.25 The reason for the negative correlation of global ecircRNA abundance with proliferation may be that ecircRNAs are more stable than their co-linear isoforms, and prefer to accumulate in cells with a slower division rate.25 A similar phenomenon was observed in fission yeast (Schizosaccharomyces pombe): after starvation, decreased cell proliferation was accompanied by an increase in ecircRNA abundance.67 In addition, accumulation of ecircRNAs is decreased in various cancer cells and idiopathic pulmonary fibrosis, as compared with that in normal tissues.25 Expression profiles of ecircRNAs were more diverse among cancer cell lines than among noncancer cells, whereas the opposite trend was observed for their corresponding co-linear isoforms.26 These observations revealed that changes in the physical condition of tissues or cells through developmental processes, aging, or disease can affect ecircRNA accumulation, suggesting that ecircRNAs might be an important biomarker for monitoring such changes. Whether (and perhaps, how) the change in ecircRNA abundance itself causes the change in cellular physical condition awaits further investigation.

Interactions with RNA-binding Proteins

It has been shown that RNA-binding proteins (RBPs), such as Argonaute,13,14 RNA polymerase II,20 and MBL,21 can bind to ecircRNAs. The protein binding capacity of ecircRNAs is likely to be more complex than previously thought. Some ecircRNAs can store, sort, or localize RBPs, and probably regulate the function of RBPs by acting as competing elements, in the same way as they modulate miRNA activity.13,68 The unique tertiary structure of ecircRNAs may also play an important role in the assembly of RNA or RBP complexes (Box 1).

CONCLUSION

It is now generally believed that eukaryotic spliceosomes exhibit a certain degree of flexibility in splice site choice, resulting in widespread alternatively
WIREs RNA Biogenesis, identification, and function of exonic circular RNAs

BOX 1

MAMMALIAN CIRCULAR INTRONIC RNAs

Through the spliceosomal machinery, intron lariats can escape the usual intron debranching and degeneration processes, and thus form stable circular intronic RNAs (ciRNAs).20 The formation process was suggested to rely on a 7 nt GU-rich motif near the 5′ splice site and a 11 nt C-rich motif near the branchpoint site.20 Although ciRNAs were first observed in mammalian cells over two decades ago,69,70 the function of ciRNAs is understudied. Recently, a comprehensive analysis of RNA-seq data in human cells provided several clues to their function.20 First, ciRNAs tend to be enriched in the nucleus and do not exhibit an excess of microRNA target sites. Second, some ciRNAs (e.g., ci-ankrd52, ci-mcm5, and ci-sirt7) can enhance expression of their parent mRNAs. Third, ciRNAs can interact with RNA polymerase II and regulate polymerase II transcription. Fourth, in some cases, the expression of ciRNAs is positively correlated with that of their parent genes. Fifth, ciRNAs tend to exhibit relatively little evolutionary conservation between human and mouse. These observations indicate that ciRNA may possess a biological role distinct from that of ecircRNAs. In addition, ciRNA may associate with the polymerase II elongation machinery and upregulate their corresponding parent genes.20 The low evolutionary conservation of intronic sequences further suggests that ciRNAs increase transcriptome complexity between species.

Biogenesis

- RCS? Regulatory factor?
- Longer intron? Expression level
- Co-/post transcription? Tissue/cell type specificity?

Function

- Translation? Conservation?
- Different types of NCL event? Localization?

Identification

- Sequence context? Sequencing error?
- Conservation? Alignment error?
- In vitro artifact?

FIGURE 6 | Summary of selected unanswered questions regarding ecircRNA biogenesis, function, and identification.

spliced isoforms, including circRNAs, in transcriptomes. Although the functions of ecircRNAs are mostly unknown, circRNAs tend to exhibit tissue/cell type-specific accumulation, and some of them are conserved among species. In addition to their well-documented role as microRNA sponges, some evidence shows that they may regulate their parental gene transcription and cell proliferation. Thus, their existence cannot be simply explained as a consequence of missplicing or an inconsequential by-product of pre-mRNA splicing. Some ecircRNAs are indeed functional. However, a lot of questions regarding ecircRNA biogenesis and degradation await answers. For example, base-pairing of matched RCSs in the flanking introns of circles is important, but not necessary, for enhancing ecircRNA formation.30,31 It is worth asking whether this property of RCSs is common to all species, and whether ecircRNA biogenesis varies among species. In addition, ecircRNAs tend to have longer flanking introns,11,15,24,37 but there is no information about how longer flanking introns are related to ecircRNA formation. Moreover, binding between splicing factor MBL proteins and MBL motifs located in flanking introns may promote ecircRNA formation from MBL genes.21 This finding has drawn great attention as MBL is known to be important in tissue-specific alternative splicing.71 Besides MBL, however, no other trans-splicing element affecting backsplicing has been found. To date, studies of circRNAs have mainly focused on ecircRNAs formed by spliceosomal machinery. Little is known about circRNAs generated from non-spliceosomal mechanisms. Their prevalence and whether they carry functions similar to those formed by spliceosomes are still awaiting further investigation. In addition, whether circRNAs are generated co-transcriptionally or post-transcriptionally remains a matter of debate. While co-transcriptional generation of ecircRNAs is supported by fly nascent RNA-seq data,21 (1) the requirement for a downstream functional 3′ processing signal and (2) the collaboration between intronic repeats and exonic sequences in circularization suggest a posttranscriptional model.32

In term of ecircRNA identification, RNA-seq data continue to be important sources, as the accessibility of such data increases exponentially with the use of different experimental designs and variously processed biosamples. The three major hurdles to ecircRNA detection from RNA-seq data are as follows: (1) discrimination between ecircRNAs, trans-spliced
RNAs, and genetic rearrangements; (2) removal of sequencing errors, alignment errors, and in vitro artifacts; and (3) biased identification results due to the use of different bioinformatics methods or sequencing data derived from different treatments. These difficulties may introduce severe bias into the trends of ecircRNA analysis. To date, there is no systematic method to effectively distinguish between different types of NCL events (i.e., circRNAs, trans-spliced RNAs, and genetic rearrangements). In addition, with currently available bioinformatics algorithms, it remains a considerable challenge to effectively eliminate false calls from sequencing/alignment errors without losing sensitivity. Our results showed that the identification results vary dramatically among different ecircRNA-detecting tools and among different RNA-treatment data (Figure 5(a)); in particular, the sensitivity varies considerably for lowly expressed circRNAs (Figure 5(b)). Our evaluations of the accuracy for five well-known circRNA-detecting methods (TopHat-Fusion, MapSplice, segemehl, find_circ, and CIRI) revealed that TopHap-Fusion yielded the best sensitivity but the worst precision, whereas MapSplice demonstrated the opposite trend (Figure 5(b)). Of these five methods, segemehl seemed to achieve the greatest balance between sensitivity and precision (Figure 5(b)). Our observation reveals that there remains a need for a robust pipeline capable of identifying ecircRNAs with better balance between sensitivity and precision. In addition, the frequent occurrence of template switching in cDNA products presents another challenge to the accurate identification of ecircRNAs. Currently, no systematic approach is available for identifying ecircRNAs in the human transcriptome while using control experiments to remove potential template switching events. In summary, there are still a lot of unanswered questions for this important but largely uncharted class of transcripts, including unknown factors relating to ecircRNA biogenesis, function, and identification (Figure 6). The world of the transcriptome may be more complicated than we previously thought.

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FURTHER READING

CircBase, a database for circular RNAs, http://circbase.org/.
circ2Traits, a database of human circular RNAs associated with disease or traits, http://gyanxet-beta.com/circdb/.
TopHat-Fusion (version 2.0.13), an algorithm for detecting both intergenic and intragenic NCL events, http://ccb.jhu.edu/software/tophat/fusion_index.html.
MapSplice (version 2.17), an algorithm for detecting both intergenic and intragenic NCL events, http://www.netlab.uky.edu/p/bioinfo/MapSplice2.
segemehl (version 0.2.0), an algorithm for detecting both intergenic and intragenic NCL events (filterjunctions.py was applied to extracting circRNA candidates), http://www.bioinf.uni-leipzig.de/Software/segemehl/.
find_circ (version 1.0), an algorithm for detecting circRNAs, https://github.com/bioxfu/circRNAFinder/blob/master/src/find_circ.py.
CIRI (version 1.1), an algorithm for detecting circRNAs, http://sourceforge.net/projects/ciri/.
Simulated datasets, the simulated datasets that were generated in this study for evaluation of the sensitivity and precision of the circRNA-detecting tools examined, ftp://treeslab1.genomics.sinica.edu.tw/simulated_expression/.

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