Circular RNAs: a new frontier in the study of human diseases

Yonghua Chen,1 Cheng Li,2 Chunlu Tan,1 Xubao Liu1

ABSTRACT
Circular RNAs (circRNAs) are recently discovered new endogenous non-coding RNAs and an area of much research activity. In addition to their potential as major gene regulators, reports are listing heterogeneous circRNA groups with many different human disorders, especially cancer. In this review, we focus on the rapidly advancing field of circRNAs that play a part in human diseases. We list tools (eg, public databases) that scan genome spans of interest to identify known circRNAs; describe the relationship between dysregulated circRNAs and human disease, highlighting their specific roles; and consider the possible use of current and potential circRNA research applications in treating human diseases. Specifically, we review the role of circRNAs as biomarkers, drug targets and therapeutic agents.

INTRODUCTION
Circular RNAs (circRNAs) are recently discovered new endogenous non-coding RNA (ncRNA) and an area of much research activity.1–3 CircRNAs are a large class of RNAs expressed in a tissue-specific and developmental stage-specific manner.1 In addition to their potential as a major gene regulator, the heterogeneous group of circRNAs may contribute to the development of many different human disorders.4 5 CircRNAs are unusually stable RNA molecules, presumably because their lack of an open end prevents conventional RNA degradation pathways. Thus, circRNAs can indicate gene expression patterns and may be an interesting, new class of biomarkers.

In this review, we focus on circRNAs involved in disease. First, we discuss tools (eg, public databases) that scan genome spans of interest to identify known circRNAs. Some of the databases search for circRNAs that are involved in a specific process or disease (eg, cancer). Second, we present disease-oriented circRNAs and discuss their relationship with the disease. Finally, we consider their possible clinical implications. Specifically, we review circRNAs as biomarkers, drug targets and therapeutic agents.

CLASSIFICATION OF CIRCRRNAS AND PUBLIC DATABASES
Recently, thousands of human circRNAs were identified using molecular biology strategies coupled with new bioinformatic approaches. Generating comprehensive circRNA classifications is not an easy task. Some circRNAs are only described in one published study. The same circRNA may be listed under different groups, depending on the classification system. For example, circRNAs predicted by computational models are often listed under different names compared with databases obtained from sequencing projects. Our understanding of circRNA biological characteristics and functions is limited; therefore, classifying circRNAs can be achieved only by focusing on their components. The following three types of circRNAs are discussed: intronic circRNAs, exonic circRNAs and exon–intron circRNAs. Intronic circRNAs are produced by connecting two or more introns, which is rare in eukaryotic cells.6 Exonic circRNAs are large molecules comprising exons and are considered by-products of exon skipping, either in pre-messenger RNA (mRNA) splicing or in mature mRNA re-splicing. The exon–intron circRNAs are enriched at transcription sites and may promote transcription of their parent mRNAs.7 8

To enable organisation of circRNAs, we provide the current online databases (table 1). These databases collect circRNAs from GenBank annotations or published articles. They list ncRNAs that have been experimentally proved, those that are purely computational predictions and those annotated as ncRNAs based on the open reading frame predicted size.

Both starBase v2.0 and circBase allow the user to search for functional classes or processes; Circ2Traits10 and nc2Cancer11 allow the user to search also by disease (eg, cancer). Although the expression datasets are not cancer-oriented, we predict a merging of the circRNA expression data bases, listed in table 1 and other datasets that are more disease-oriented (eg, Circ2Traits; http://gyanxet-beta.com/circdb/). At this time, the genomic positions of several circRNAs can be matched to databases that annotate SNPs associated with disease or disease-associated genetic regions (Circ2Traits10 and nc2Cancer11). Ghosal et al10 measured the likelihood of disease association for a given circRNA from the statistical significance of the interaction of circRNAs with the disease-associated microRNAs (miRNAs). They then analysed gene ontology enrichment on the protein coding genes in the miRNA–circRNA disease interactome to identify enriched genes associated with particular biological processes. Biological process enrichment for mRNAs in 90 diseases was identified. Among these mRNAs, there are 22 light stimulus response genes and 43 cell cycle–related genes associated with breast cancer.10 This is the first comprehensive data analysis investigating the global effects of the potential association between circRNAs and cancer.

Recent studies show that miRNAs interact with ncRNAs, such as circRNAs.3 11 The circRNAs and mRNAs with common miRNA target sites compete for miRNA binding and form a complex network.
CircRNAs participate in a wide range of biological processes. Almost every step in a gene’s life cycle, including transcription, mRNA splicing, RNA decay and translation, can be influenced by circRNAs. Considering the wide range of roles that circRNAs play in cellular networks, it is not surprising that their misregulation leads to abnormal cellular functions and growth defects and is implicated in human disease. We focused on developing a list of circRNAs linked to human disease by various means. We mainly used two of the online databases to retrieve circRNAs (Circ2Traits10 and nc2Cancer11 databases) and we searched PubMed for articles linking these circRNAs to human disease. In table 2, we summarise our findings.

Several circRNAs are causally involved in human disease. In some cases, the link between the circRNA and human disease was obvious and human disease was the model in which these circRNAs were first described (eg, the exonic circRNAs that are antisense cerebellar degeneration-related protein 1 transcript (CDR1as) and sex-determining region Y (Sry)). However, we also found some circRNAs with incompletely elucidated links to human disease, but preliminary findings suggest that further investigation of possible connections is needed. Here, we focus on the genetic and epigenetic events that disrupt circRNA loci and related proteins in cancer and other human diseases, such as cancer pathogenesis. Mutations in miRNAs (especially in seed regions) and their target sites may alter miRNA–circRNA interactions and rewire the ceRNA network. The SomamiR 2.0 database contains somatic mutations in miRNAs and their target sites for three types of ceRNAs—circRNAs, long ncRNAs and miRNAs.20

### CIRCNRNAS IN HUMAN DISEASES

Table 1 Public circRNA databases

<table>
<thead>
<tr>
<th>Name</th>
<th>circ2Traits10</th>
<th>nc2Cancer11</th>
<th>circBase12</th>
<th>starBase v2.013</th>
<th>CircNet14</th>
<th>deepBase v2.015</th>
<th>CircInteractome16</th>
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<tr>
<td>CircRNA disease association</td>
<td>105 Diseases</td>
<td>31 Cancers</td>
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<td>Not available</td>
<td>Not available</td>
<td>Not available</td>
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<td>172 Human circRNAs</td>
<td>Not available</td>
<td>Not available</td>
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<td>212 950 CircRNAs</td>
<td>14 867 Human circRNAs</td>
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<tr>
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<td>Not available</td>
<td>A web interface of BLAST</td>
<td>Not available</td>
<td>A web interface of BLAST</td>
<td>Not available</td>
<td>Not available</td>
</tr>
<tr>
<td>CircRNA reference source</td>
<td>circRNA dataset from Memczak et al1</td>
<td>miR2Disease, miRCancer, HMDD v2.0 and starBase</td>
<td>Back-spliced junction sites in animals reported in 2013 and 2015</td>
<td>circBase v0.1</td>
<td>Reported human circRNAs from 2013 to 2015</td>
<td>NCBI GEO and SRA databases</td>
<td>CircBase, starBase V.2.0, miRBase,17 IRESite18</td>
</tr>
<tr>
<td>CircRNA position on genome</td>
<td>A customised genome browser</td>
<td>Not available</td>
<td>Linked out to UCSC Genome Browser</td>
<td>A customised genome browser accessible through keyword search</td>
<td>An integrated genome browser synchronising with the network graphical user interface</td>
<td>A customised genome browser</td>
<td>Linked out to UCSC genome browser</td>
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<tr>
<td>CircRNA sample source</td>
<td>Not available</td>
<td>Not available</td>
<td>The samples where the back-spliced junction sites were discovered</td>
<td>circBase v0.1 source samples</td>
<td>1. In which sample junction sites were discovered. 2. Expression level in available samples. 3. Clustered sample conditions</td>
<td>Not available</td>
<td>Not available</td>
</tr>
<tr>
<td>CircRNA naming</td>
<td>A serial number for every detected back-spliced junction site</td>
<td>A serial number for every detected back-spliced junction site</td>
<td>A serial number for every detected back-spliced junction site</td>
<td>Same as circBase, except CDRI antisense (CDR1as)</td>
<td>A systematic naming system which provides information to the source gene and annotated exons of circRNAs</td>
<td>A systematic naming system which provides information to the transcript number</td>
<td>Same as circBase</td>
</tr>
<tr>
<td>CircRNA expression profiles in samples</td>
<td>Not available</td>
<td>Not available</td>
<td>Not available</td>
<td>Not available</td>
<td>An all-sample expression heat-map for every circRNA and linear isoform</td>
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<tr>
<td>Address on miRNA regulatory relationships</td>
<td>Identifies circRNA and miRNA interactions</td>
<td>Identifies circRNA and miRNA interactions</td>
<td>Identifies circRNA and miRNA interactions through Chip-Seq data analysis</td>
<td>A network-driven graphical interface shows the relationship between miRNA target genes and circRNAs</td>
<td>Not available</td>
<td>Identifies circRNA and miRNA interactions</td>
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<td>RNA-binding protein</td>
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<td>Not available</td>
<td>Not available</td>
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<tr>
<td>circRNA isoforms</td>
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<td>Not available</td>
<td>Not available</td>
<td>Not available</td>
<td>All traceable on the integrated genome browser</td>
<td>All traceable on the integrated genome browser</td>
<td>Not available</td>
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</table>

of interaction and regulation, commonly known as the competing endogenous RNA (ceRNA) network. Dysregulated crosstalk between ceRNAs in the network plays an important role in cancer pathogenesis. Mutations in miRNAs (especially in seed regions) and their target sites may alter miRNA–ceRNA interactions and rewire the ceRNA network. The SomamiR 2.0 database contains somatic mutations in miRNAs and their target sites for three types of ceRNAs—circRNAs, long ncRNAs and miRNAs.20
as neurological, cardiovascular and autoimmune disorders. We next illustrate several circRNAs and their involvement in human diseases.

**CircRNAs: deregulation in cancer**

In cancer biology, the search for gene expression differences between tumour and normal samples is considered most important, as knowledge of circRNA expression profiles in tumour and normal samples is modest. Commercial gene expression arrays used for polycomb group proteins may contain probes that hybridise to circRNAs and it may be possible to retrieve cancer-related circRNA expression profiles from public, tumour-specific, gene-expression datasets (eg, Circ2Traits and nc2Cancer). To identify new transcripts, some investigators have used the Arraystar Human CircRNA array, which can test for circRNA gene expression.

Table 2  CircRNAs associated with human disease

<table>
<thead>
<tr>
<th>CircRNA</th>
<th>Alias</th>
<th>Regulation</th>
<th>Gene symbol</th>
<th>Sample</th>
<th>Disease/tumour</th>
<th>Genome position</th>
</tr>
</thead>
<tbody>
<tr>
<td>hsa_circ_0001946</td>
<td>CDR1as</td>
<td>–</td>
<td>CDR1</td>
<td>Cell lines</td>
<td>Colorectal cancer,(^{22}) breast cancer(^{22})</td>
<td>chr13:139865339–139866824</td>
</tr>
<tr>
<td>hsa_circ_0001946</td>
<td>CDR1as</td>
<td>↓</td>
<td>CDR1</td>
<td>Human samples</td>
<td>Alzheimer’s disease(^{25})</td>
<td>chr13:139865339–139866824</td>
</tr>
<tr>
<td>hsa_circ_0001141</td>
<td>hsa_circ_001763</td>
<td>↓</td>
<td>ITCH</td>
<td>Human samples</td>
<td>Colorectal cancer,(^{26}) oesophageal carcinoma(^{27})</td>
<td>chr20:33001547–33037285</td>
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<tr>
<td>hsa_circ_0006229</td>
<td>–</td>
<td>↓</td>
<td>TNS3</td>
<td>Human samples</td>
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<td>AZIN1</td>
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<td>hsa_circ_0002190</td>
<td>circ-KLDHC10</td>
<td>↑</td>
<td>KLDHC10</td>
<td>Human samples (cancer serum)</td>
<td>Colorectal cancer(^{29})</td>
<td>chr7:129760588–129762042</td>
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<td>hsa_circ_0001649</td>
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<td>↓</td>
<td>SHPRH</td>
<td>Human samples</td>
<td>Hepatocellular carcinoma(^{30})</td>
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<td>hsa_circ_0000140</td>
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<td>↓</td>
<td>KIAA0907</td>
<td>Human samples</td>
<td>Gastric cancer(^{31})</td>
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<td>hsa_circ_0010783</td>
<td>–</td>
<td>↑</td>
<td>PCDH9</td>
<td>Human samples</td>
<td>Immunosenescence(^{32})</td>
<td>chr6:167750008–167750243</td>
</tr>
</tbody>
</table>

Li et al\(^{29}\) examined serum exosome (small membrane vesicles of endocytic origins secreted by cells) RNA sequencing datasets from 11 patients with CRC and normal serum. In comparison with healthy subjects, 67 circRNAs were missing and 257 new circRNA species were detected in the patients with CRC. Notably, many of the host genes for these new circRNAs (48 genes for 53 circRNAs) were significantly upregulated in CRC tissues. qRT-PCR analysis confirmed that the expression levels of circ-KLDHC10 (also known as hsa_circ_0002190) were upregulated in CRC serum\(^{29}\).

In addition, a recent review reported that expression analyses of various tumour cell lines showed widespread expression of ciRS-7 in neuroblastomas, astrocytoma, renal cell and lung carcinomas\(^{30}\). The stable expression of ciRS-7 in HeLa cells\(^{31}\) indicates that ciRS-7 may be associated with cervical cancer.

CircRNAs are differentially expressed in a wide range of cancers and may be involved in cancer initiation and progression. Continuing research into circRNAs shows that they may play roles in other tumours and are potential new biomarkers for diagnosis. For example, serum exosomal circRNAs profiles can distinguish between patients with cancer and healthy controls\(^{29}\).

CircRNAs are associated with cancer-related miRNAs

It is clear that miRNAs are involved in nearly all aspects of cellular functions\(^{32}\) and have a critical role in cancer initiation and progression\(^{33}\)\(^{36}\). In March 2013, two publications reported that circRNAs function as miRNA ‘sponges’, which naturally sequester and competitively suppress the activity of miRNAs\(^{1}\)\(^{19}\). The circRNAs and miRNAs with common miRNA target sites compete for miRNA binding and form a complex interactive and regulatory network, commonly known as the ceRNA network. The miRNA target recognition is largely dependent on sequence complementarity between the miRNA seed region (nucleotides 2–7 in the mature miRNA sequence) and its target sites on ceRNAs. Mutations in miRNAs (especially the seed regions) and their target sites may alter miRNA–ceRNA interactions and rewire the ceRNA network\(^{37}\). The dysregulation of crosstalk between ceRNAs in the network has an important role in cancer pathogenesis, suggesting that circRNAs might be involved in malignant tumours correlated with miRNAs.

By measuring the likelihood of a circRNA association with disease from the statistical significance of their interaction with miRNAs associated with the disease in question, the Circ2Traits database\(^{10}\) listed biological processes for miRNAs in 90 diseases. Among these miRNAs, there are 22 light stimulus response genes and 43 cell cycle-related genes associated with breast cancer\(^{10}\). This is the first report with a global view of the potential association between circRNAs and cancer based on comprehensive

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data analysis. However, a direct circRNA and miRNA association requires more biological evidence. The SomamIR 2.0 database contains somatic mutations in miRNAs and their target sites on circRNAs, long ncRNAs and miRNAs.20

Identifying ceRNAs and circRNAs as important regulators of miRNA activity underlines the increasing complexity of ncRNA-mediated regulatory networks. In particular, the recently identified circRNA, known as ciRS-7, which acts as a ceRNA or super sponge of miR-7, competitively inhibits miR-7 activity and promotes oncogene expression (such as EGFR and XIAP), while inhibiting tumour suppression genes (such as KLF4) and therefore promoting the initiation and development of cancer, such as HCC, breast cancer and cervical cancer.7 22 Hence, discovering the regulation of miR-7/miR-671/ciRS-7 axis activity will probably advance the understanding of various cancer aetiologies.5 Another notable circular miRNA sponge is Sry. Sry controls the biological effects of miR-138 by binding to its 16 conserved binding sites.19 It can regulate complex regulatory networks and influence many physiological and pathological processes, by modulating multiple miRNAs. Because both Sry and miR-7 have a crucial effect on the occurrence and progression of cancer, we assume that circRNAs must also be involved in the process.

CircRNA–miRNA axes regulate cancer-related pathways

Some endogenous circRNAs in mammalian cells are highly abundant and evolutionarily conserved in oncogenesis. To date, the evidence suggests that circRNAs may regulate transcription and pathways by manipulating miRNAs. The ciRS-7/miR-7 axis is probably involved in cancer-associated biological processes, such as cancer initiation and progression.3 22 The circRNA regulatory influence on miR-7 is clear; ciRS-7 overexpression acts as a miRNA sponge to restrain the expression miR-7 and therefore elevates EGFR expression. ciRS-7 can naturally sequester and inhibit miR-7 activity and promote oncogenic EGFR and XIAP gene expression as well as inhibit the tumour-suppressor KLF4 expression, thus promoting the initiation and development of cancer, such as cervical cancer, breast cancer and HCC.5 22 Additionally, miR-7 indirectly upregulates E-cadherin by targeting FAK24 and IGF1R,38 resulting in reduced epithelial to mesenchymal transition (EMT), decreased anchorage-independent growth and suppression of metastasis.3

Li et al27 have shown that circ-ITCH may have an antitumour function in oesophageal squamous cell carcinoma through interactions with miRNAs such as miR-7, miR-17 and miR-214 and an increase in ITCH. These interactions facilitate ubiquitin-mediated Dvl2 degradation and decrease expression of the oncogene c-myc. This process therefore inhibits canonical Wnt signalling. In addition, Huang et al26 drew the parallel conclusion that circ-ITCH expression is typically downregulated in CRC, and that circ-ITCH has an inhibitory role in the canonical Wnt pathway, inhibiting c-myc and cyclinD1 expression.

A recent study indicated that the majority of circRNAs are regulated by human EMT and more than one-third of abundant circRNAs are dynamically regulated by the alternative splicing factor, Quaking, which is regulated by the EMT process.39 Given that EMT participates in tumorigenesis and provides important indications for cancer progression, we may obtain more knowledge of the therapeutic role of circRNA by targeting miRNAs involved in EMT.

CircRNAs involved in neurological disorder

CircRNAs play a critical role in normal nervous system function and during various differentiation stages. Most circRNA expression is associated with specific neuroanatomical regions, cell types or subcellular compartments. Moreover, circRNA levels increase relative to linear mRNAs during ageing.40 41 Recently, two papers by Rybak-Wolf et al42 and You et al43 provide a valuable circRNA catalogue of the mammalian brain and shed a new light on their potential function in the nervous system. The authors42 43 report the identification of thousands of conserved circRNAs that are highly expressed in mammalian brain. Many of these circRNAs are upregulated during neurogenesis and further enriched in synaptic processes compared with their linear isoforms. These findings further highlight a potential role of brain circRNAs in the nervous system. Rybak-Wolf et al42 examined 29 human/mouse RNA sequencing datasets from dissected brain tissues or neuronal-differentiated cell lines. They found that some circRNAs are expressed dynamically and independently of their linear transcripts, implying regulated expression of brain circRNAs. You et al43 drew the parallel conclusion that circRNAs are highly enriched and developmentally regulated in the brain. Therefore, circRNA dysregulation causes various neurodegenerative and neurological disorders. CircRNAs have been implicated in neurological disorders, such as Alzheimer’s disease (AD),42 Parkinson’s disease10 and many others. We review the association of various circRNAs with neurological disorders.

The discovery of CDR1as, a miR-7 sponge, promoted theories about the link between circRNAs and neurodegenerative diseases such as AD, since miR-7 and some other microRNAs are directly related to neurodegenerative diseases.24 For example, CDR1as is highly expressed in the brain and has over 60 binding sites for miR-7.1 19 44 It is important to note that miR-7 is implicated in numerous pathways and diseases, including its function as a direct regulator of α-synuclein and ubiquitin protein ligase A. CDR1as has been implicated in Parkinson’s disease,10 AD and brain development.19 A recent report supports the downregulation of CDR1as in patients with Alzheimer’s disease.25 Using northern blot hybridisation techniques and the circularity-sensitive circRNA probe, RNaseR, Lukiw25 provided evidence of a misregulated miR-7–circRNA system in the sporadic AD hippocampal CA1 region. The author found that deficits in ciRS-7 and ciRS-7 ‘sponging activities’ increase ambient miR-7 levels in AD-affected brain cells, ultimately downregulating selective miR-7 targets. Upregulated miR-7, due to a deficiency in ciRS-7 ‘sponging’ effects, is predicted to downregulate AD-relevant targets, such as the ubiquitin protein ligase A (UBE2A). UBE2A, which is depleted in AD brains, is an autophagic, phagocytic protein that is essential for the clearance of amyloid peptides in AD and other progressive inflammatory degenerations of the human central nervous system.

Amyotrophic lateral sclerosis is a devastating neurodegenerative disease, primarily affecting motor neurons. Mutations in the gene encoding TDP-43 cause some forms of the disease, and cytoplasmic TDP-43 aggregates accumulate in degenerating neurons of most people with amyotrophic lateral sclerosis. Thus, strategies targeting the toxicity of cytoplasmic TDP-43 aggregates may be effective. A study by Armakola et al45 shows that in the absence of Dbr1 enzymatic activity, intronic lariats accumulate in the cytoplasm to sequester TDP-43, preventing the aggregates from interfering with essential cellular RNAs and RNA-binding proteins. Dbr1 knockdown in a human neuronal cell line or in primary rat neurons is sufficient to suppress TDP-43 toxicity.46 47

These studies suggest that circRNAs have a crucial role in neurological functions, although their mechanisms are unknown. The specific expression and stability of circRNAs...
make them interesting candidates as biomarkers for neurodegenerative diseases, such as AD.

Disruption of circRNAs in other diseases
CircRNAs are disrupted and associated with cardiovascular disorders. For atherosclerotic vascular disease (ASVD), Burd et al. discovered that expression of the new circRNA transcript circular antisense non-coding RNA in the INK4 locus (cANRIL) may be correlated with INK4/ARF transcription and ASVD risk. Based on genome-wide association study studies, SNPs in non-coding regions have been shown to be associated with higher susceptibility to a range of diseases. A large number of SNPs in the INK4/ARF loci were associated with increased ASVD risk. The chromosome region in which the SNPs were characterised harbours the protein-coding genes CDKN2a (INK4a) and CDKN2b (INK4b). Both these genes are adjacent to the gene encoding cANRIL. cANRIL is an antisense transcript from the INK4A/ARF locus. SNPs on chromosome 9p21.3 near the INK4/ARF (CDKN2a/b) locus within the ASVD risk interval may modulate ANRIL splicing and cANRIL production. Hence, cANRIL could be involved in ASVD.

Ashwal-Fluss et al. discovered that a new circRNA transcript called CircMbl might be correlated with myotonic dystrophy initiation and progression. CircMbl and its flanking intron sequences can combine with MBL. Alterations in MBL levels strongly affect circMbl biosynthesis. CircRNA production competes with canonical mbl pre-mRNA splicing. MBL can regulate mbl pre-mRNA splicing efficiency through the activity of both mbl mRNA and circMbl. Moreover, circMbl can act as an MBL sponge. MBL functional deficiency causes a severe degenerative disease called myotonic dystrophy. Hence, circMbl could be involved in myotonic dystrophy initiation and progression.

Aberrant immune responses are a hallmark of ageing and age-associated diseases, associated, in part, with the continuous proinflammatory cytokine secretion by senescent cells. Recent findings link circRNAs to CD28-related CD8(+) T cell ageing and global immunosenescence. Using cross-comparison of circRNA microarrays and stepwise bioinformatic assays, Wang et al. investigated circular RNA–micro RNA interactions in ageing human CD8(+) T cell populations and examined the accompanying loss of CD28 expression. The authors discovered that circRNA100783 might regulate phosphoprotein-related signal transduction during CD28-dependent CD8(+) T cell ageing.

The current circRNA field is based upon disease-associated changes in circRNA expression. However, genetic studies on circRNA sequences may distinguish the specific contribution of large- and small-scale mutations to circRNA functions. With improved understanding of circRNA language, we will be able to classify diseases based on the mutations identified and their effect on circRNA function.

Diagnostic and prognostic biomarkers
According to the current literature, the main characteristics of circRNAs are as follows: (i) universality; (ii) conservatism, as the signal behind circularisation seems to be evolutionarily conserved among different species; (iii) definite specificity; (iv) stability, or a resistance to debranching enzymes and RNA exonucleases and (v) highly abundant expression, as the gene product level of some exonic circRNAs is higher than that of linear RNAs. Therefore, circRNAs possess distinct advantages and may be useful as new biomarkers for diagnosis, prognosis and therapeutic response prediction.

The marked increase or decrease in circRNA expression levels in tumours compared with normal tissues seems to be a feature shared by circRNAs that would be useful in diagnostics. There are a few examples of circRNAs with a diagnostic role. For example, several circRNAs are reportedly aberrantly expressed in human cancers (such as, oesophageal carcinoma, CRC and gastric cancer). Upregulation or downregulation of circRNAs in cancer tissues compared with paired adjacent tissue could indicate a diagnostic potential new biomarker. A large difference in circRNA tumour expression levels compared with normal tissues is a topic for future clinical research, although larger clinical datasets need to be assayed. Other circRNAs might also be promising biomarkers. CircRNAs have been described as an ageing biomarker class in Drosophila.

A potential avenue of circRNA research is the circulating circRNAs in serum, plasma and other body fluids, especially in microparticles, like exosomes. One of the main advantages of circRNAs is their high circulating stability. They can be detected through minimally invasive blood, urine or saliva sample collection coupled with RT-PCR. This might represent an unexpected and unexplored potential disease biomarker for diagnosis, prognosis and therapeutic response prediction.

Contribution to targeted therapy
CircRNAs might be useful therapeutic agents. Circularisation may be a future target for treatment, either to reduce the circularisation of functional transcripts or to use an ‘mRNA trap’ to sequester dysfunctional exons in transcripts. Circularising the miRNA sponge in cells is a new candidate for RNA-based cancer treatments. Researchers recently reported that a circular, artificial, miRNA sponge might induce miRNA loss-of-function in cancer cells. Liu et al. constructed a circular miRNA sponge expression vector containing the permuted intron–exon sequence derived from the group I intron of T4 bacteriophage gene td and produced circular miRNA sponges against miR-21 and miR-221. As alternative vectors expressing linear sponges, the expression vectors for RNA circles described in this study open new ways to deliver miRNA sponges with persistent effects and hold great potential for cancer research and treatment. Endogenous circRNAs were recently identified as a new class of gene expression regulators, acting as miRNA sponges in mammals. ciRS-7 functions as a potent circular miR-7 sponge, containing multiple miRNA response elements that bind miR-7. Thus, it can instantaneously bind or release a large number of miR-7 molecules, thereby effectively regulating the disease network. The miRNA sponge function of endogenous circRNAs is a general phenomenon. Studying the evolutionarily optimised, circular miRNA sponge structures should provide valuable insights into the design and development of potent artificial sponges in order to achieve effective inhibition of miRNA function in diseases.

ncRNAs broaden the universe of potential ‘druggable’ targets. The scientific community and pharmaceutical companies must pursue these new approaches vigorously, using automated large-scale screening of these miRNA-related drugs, developing knock-in and knockout models for the target circRNAs, etc. Targeting circRNAs and human diseases, in addition to miRNAs, is still in its infancy but important developments are expected.

FUTURE PERSPECTIVES

Interest in the contribution of circRNAs to the genesis and progression of human disorders is growing. While new information and insights into circular RNAs are generated rapidly, the biological and molecular mechanisms of circRNAs in the development of diverse diseases are not yet fully understood. These are likely to include miRNA sponging, splicing regulation and safeguarding for the assembly of macromolecular complexes (such as circRNA–protein complexes). In particular, little is known of the molecular mechanisms of their biogenesis, degradation and cellular localization. It should be borne in mind that circRNAs may be a large family with widely diverse biogenesis, degradation, cellular localization, function and molecular mechanisms. Additional circRNAs will be identified as technology and research develops. Moreover, functional studies will disclose both physiological and pathological processes of the vast majority of circRNAs. We predict the construction of engineered circRNAs as molecular tools or treatments. Engineered circRNAs might be effective either for sequestering many RNAs and RNA-binding proteins or for releasing these stored molecules via cleavage of the circRNA.

CircRNAs provide new insights into the ‘dark matter’ of the human genome. The research and application prospects for circRNAs in human disorders are promising. CircRNAs may affect life processes, serve as diagnostic or predictive biomarkers of disease and provide new potential therapeutic targets. Our hope is that chemical and biotechnological advances will take place alongside basic studies, disclosing the physiological and pathological functions of circRNAs and developing circRNA-based therapeutic strategies, allowing safe and successful inclusion in day-to-day clinical practice.

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The authors apologise to those whose work has not been cited owing to limitations of space.

Contributors

C-YH and L-C contributed equally to this paper; C-YH and L-C read and analysed the relevant literature; C-YH, L-C and T-CL wrote the manuscript; L-XB revised it.

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