Closing the circle: current state and perspectives of circular RNA databases

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Abstract
Circular RNAs (circRNAs) are covalently closed RNA molecules that have been linked to various diseases, including cancer. However, a precise function and working mechanism are lacking for the larger majority. Following many different experimental and computational approaches to identify circRNAs, multiple circRNA databases were developed as well. Unfortunately, there are several major issues with the current circRNA databases, which substantially hamper progression in the field. First, as the overlap in content is limited, a true reference set of circRNAs is lacking. This results from the low abundance and highly specific expression of circRNAs, and varying sequencing methods, data-analysis pipelines, and circRNA detection tools. A second major issue is the use of ambiguous nomenclature. Thus, redundant or even conflicting names for circRNAs across different databases contribute to the reproducibility crisis. Third, circRNA databases, in essence, rely on the position of the circRNA back-splice junction, whereas alternative splicing could result in circRNAs with different length and sequence. To uniquely identify a circRNA molecule, the full circular sequence is required. Fourth, circRNA databases annotate circRNAs’ microRNA binding and protein-coding potential, but these annotations are generally based on presumed circRNA sequences. Finally, several databases are not regularly updated, contain incomplete data or suffer from connectivity issues. In this review, we present a comprehensive overview of the current circRNA databases and their content, features, and usability. In addition to discussing the current issues regarding circRNA databases, we come with important suggestions to streamline further research in this growing field.

Key words: circRNA; circRNA databases

Introduction
After their discovery more than three decades ago, circular RNAs (circRNAs) have been emerging as a large class of generally non-coding RNAs. Originating from the same precursor as linear RNA transcripts, circRNAs are formed through a process called back-splicing, in contrast to regular forward splicing. Back-splicing results in a covalently closed loop characterized by a non-linear back-spliced junction (BSJ) between a splice donor and an upstream splice acceptor, and lacking a poly(A) tail and 5' and 3' ends. Due to their circular nature, circRNAs are more resistant to degradation by exonucleases and therefore, more stable than linear RNA [1,2]. CircRNAs are widespread and abundant in a variety of organisms. It is estimated that the total number of circRNA molecules is roughly 1% of the number of poly(A) molecules [3]. Generally, the expression levels of most circRNAs is estimated to be 5–10% of their corresponding linear RNA product [4]. Interestingly, the majority of circRNAs seem to be cell-type specific [3,5].

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Although the function of most circRNA remains largely unknown, increasing evidence shows that circRNAs can act as a sponge for microRNAs (miRNA) and RNA binding proteins (RBPs), as modulator of transcription and splicing, and as template for translation [6,7]. Furthermore, circRNAs have been associated with a broad range of diseases, including various types of cancer [8,9]. Here, circRNAs have been found to act as miRNA sponges to inhibit their regulation of downstream cancer target genes. CircCDS1 and circMT01, for example, bind to miR-7 and miR-9, respectively, and influence gene regulation, thus indirectly achieving either tumor inhibition or stimulation [10,11]. Due to the observed associations between circRNA abundance and cancer, circRNAs may serve as cancer biomarkers with good diagnostic performance [12].

Various studies also demonstrated that circRNAs are present at relatively high steady state levels in human biofluids, such as saliva, plasma, serum and in exosomes, which makes them attractive candidate biomarkers for noninvasive liquid biopsies [1]. For example, circ-ZEB1.33 was overexpressed in hepatocellular carcinoma (HCC) compared to adjacent normal tissue and normal liver. In line with this, the serum level of circ-ZEB1.33 was higher in HCC patients compared to healthy controls, and its levels in HCC tissue and serum were correlated across different TNM stages (TNM Classification of Malignant Tumors) and were associated with overall survival in HCC patients [13].

Numerous bioinformatics pipelines have been developed to identify circRNAs [14–17], leading to the prediction of millions of circRNAs in different short read RNA-sequencing (RNA-seq) datasets [2,3,18,19]. This spurred in the development of more than 20 databases containing human circRNAs. These databases also contain various circRNA annotations, such as circRNA tissue- and disease-specificity, circRNA–miRNA interactions, circRNA–RBP interactions, circRNA coding potential and conservation amongst species. Each has its unique aspects and merits, but we are far off a uniform consensus circRNA catalog. In this review, we present a comprehensive overview of the current circRNA databases and their content, features and usability. Furthermore, we discuss the current issues regarding circRNA databases and come with important suggestions to streamline further research in this growing field.

Material and methods

Literature search

PubMed and Google were queried with the following keywords: ‘circRNA database’, and all relevant hits were inspected manually. To keep the focus of our analyses on circRNA databases, only databases specific for circRNAs were included in the result tables. Other databases with interesting features are mentioned in the text. Databases exclusively containing plant circRNAs [20] were not included in this review.

Data acquisition

All circRNA database websites were visited on 03 September 2019 using Google Chrome, Firefox and Safari. When available, database exports were downloaded.

Data processing

Database content was processed in RStudio (v1.2.1335). All databases containing circRNA coordinates based on the hg38 genome build were converted to hg19 using LiftOver (UCSC Genome Browser [21]). We noticed that the start positions in the files obtained from circAtlas v2.0 were one nucleotide off compared to the other databases. To compensate for this issue, the start of each BSJ in circAtlas v2.0 was lowered by one nucleotide.

The number of unique circRNAs in each database was calculated based on the BSJ or based on the unique name for the noncurated and curated databases, respectively.

Euler plots were generated with CRAN package Eulerr (v5.1.0). It is important to note that while Euler plots are a very helpful visualisation, there is always some error, and the higher the number of diagrams, the higher the error. To ensure correct interpretation of the plots, all Euler plot results are also reported. The overlap between circRNA databases was calculated based on the BSJ position. As not all databases report the strand from which the circRNA originates, circRNAs were compared solely on their BSJ position.

The number of single-exon circRNAs was calculated by comparing the BSJ positions with all exon positions (downloaded from Ensembl, GRCh37 archive [22]).

Results

Overview of human circRNA databases

In total, we selected 20 human circRNA databases and divided them into two categories: noncurated databases, based on in-house or publicly available RNA-seq or circRNA datasets; and curated databases, based on literature searches for empirically validated circRNA (Tables 1 and 2; Supplemental Table 1). Despite several attempts, Circ2Traits, CircInteractome, CircNet and deepBase v2.0 were unreachable, and therefore not included in some of our analyses. In addition, Circ2Disease and circRNADb were often found to be unreachable.

All curated circRNA databases employ the same content search strategy, namely a literature search with keywords such as ‘circular RNA’ and ‘circRNA disease’, followed by manual selection of suitable articles. Interestingly, apart from circRNA validation, Circ2Disease also includes manually curated circRNA–miRNA interactions, circRNA–RBP interactions and other up- or down-stream regulatory genes.

While most circRNA databases predominantly store human circRNAs, a few databases also include other species such as fly, worm and mouse (Figure 1). TSCD claims to contain circRNAs detected in macaque samples but these data are currently not present in the database. Interestingly, the number of circRNAs varies substantially across the databases. In addition, each circRNA database provides different types of circRNA annotations, which are described in the following paragraphs.

CircRNA annotation

To facilitate functional exploration of circRNAs, circRNA databases typically include several annotation levels. This section contains a short description of these annotations; a complete overview of circRNA annotations can be found in Supplemental Table 1.

Tissue-specificity

CircRNAs are generally annotated as tissue-specific if they are detected in specific tissues or cell types, sometimes assessed by a specificity score. CircAtlas v2.0, CIRCpedia v2 and TCSD all
## Table 1. Characteristics of noncurated circRNA databases. (A) Noncurated databases that were online accessible (B) Noncurated databases that were not online accessible

| Database          | Dataset                  | # circRNAs* | Detection tool(s) | Filter | Short description                                                                 |
|-------------------|--------------------------|-------------|-------------------|--------|-----------------------------------------------------------------------------------|
| **A**             |                          |             |                   |        |                                                                                   |
| circAtlas v2.0 [23] | In-house data            | 610,406     | CIRI2, DCC, MapSplice, CircExplorer2 | ≥2 tools ≥2 BSJ reads | Detection of circRNAs in 17 human tissues reports conservation across species, circRNA-microRNA interaction, predicted ORF, IRES and RBP sites |
| circbank [24]     | Public circRNA dataset (circBase) | 140,725   | Does not apply    | Unknown | Focus on nomenclature system; reports miRNAs binding sites, conservation across species, m6A modification, mutations in circRNA, predicted ORFs and IRES sites |
| circBase [25]     | Public circRNA datasets  | 92,375      | Depends on source | None   | Unified database of all circRNAs predicted by nine large-scale studies; reports putative spliced circRNA sequences |
| CIRCpedia v2 [26] | Public RNA-seq datasets  | 177,456     | CIRCexplorer2, MapSplice CIRI2, find_circ, CircExplorer2, circRNA_finder | ≥2 tools ≥2BSJ reads | Focus on circRNAs in 65 human cell lines |
| CircRiC [5]       | Public RNA-seq datasets  | 92,599      | CIRI2, find_circ, CircExplorer2, circRNA_finder | ≥2 tools ≥2BSJ reads | Focus on detection of lineage-specific circRNAs in 935 cancer cell lines; reports drug response, biogenesis, interactions between circRNAs and mRNA (including miRNAs), proteins, or mutations |
| circRNADb [27]    | Public circRNA datasets  | 32,914      | Depends on source | ≥2 BSJ reads | Focus on protein-coding annotation (predicted ORFs with protein-coding potential and evidence by mass spectrometry; reports exon splicing information and predicted IRE sites |
| CSCD [11]         | Public RNA-seq datasets  | 1,223,114   | CIRI2, find_circ, circRNA_finder, CircExplorer | None   | Focus on distinguishing cancer-specific circRNAs from ‘normal’ circRNAs in 87 cancer cell lines and 141 normal cell lines; reports circRNA-miRNA interactions, putative splicing possibilities, predicted cellular location, RBP sites and ORFs |
| exoRBase [28]     | Public blood exosomal RNA-seq datasets | 57,412 | ACTS, find_circ | ≥2 tools | circRNA detection in 2093 clinical human cancer samples using exome capture sequencing |
| MiOncoCirc v2.0 [8] | In-house data            | 227,056     | CIRI2explorer     | ≥2 BSJ reads | Tissue-specific circRNAs in 16 adult human tissues and 15 fetal human tissues; reports circRNA-miRNA interactions, conservation across species and predicted RBP sites |
| TSCD [29]         | Public RNA-seq datasets  | 284,296     | CIRI, find_circ, circRNAfinder | ≥2 BSJ reads | Tissue-specific circRNAs in 16 adult human tissues and 15 fetal human tissues; reports circRNA-miRNA interactions, conservation across species and predicted RBP sites |
| **B**             |                          |             |                   |        |                                                                                   |
| Circ2 Traits [30] | Public circRNA dataset   | 1953**      | find_circ         | ≥2 BSJ reads | Association of circRNAs with diseases based on the interactions of circRNAs with disease-associated miRNAs and disease-associated SNPs mapped on circRNA loci |
| CircInteractome [31] | Public circRNA dataset (circBase) | Unknown | Does not apply | Unknown | Focus on interaction between miRNAs and circRNAs with RBP sites; reports IRES sites and ORFs |
| CircNet [32]      | Public RNA-seq datasets  | 34,000**    | find_circ         | ≥3 sources | Maps circRNA-miRNA-mRNA interactions into regulatory networks |
| deepBase v2.0 [33] | Public RNA-seq and circRNA datasets (circBase) | 14,867** | find_circ | Unknown | Comparison of (small) noncoding RNAs (including circRNAs) across 19 species; reports conservation between species |

*Number of unique human circRNAs in the download files.

**Number of human circRNAs reported by the authors; this could not be verified as the database was not online.
report circRNA expression levels across various human tissues and cell lines. In addition, CircRiC focuses on circRNAs in cancer cell lines and MiOncoCirc v2.0 on circRNAs in clinical human cancer samples.

**CircRNA-disease associations**
Due to their potential use as biomarkers, there has also been increasing interest in the association of circRNAs with diseases. These associations are mostly reported by curated databases, where circRNAs are considered disease-specific when up- or downregulated in a particular disease sample. circRNADb is a noncurated circRNA database that also reports circRNA-disease associations if there is a link between the parental gene of the circRNA and a specific disease.

**CircRNA–miRNA interactions**
In total, 12 of the databases discussed in this review (60%) report circRNA–miRNA interactions. To predict miRNA binding sites in circRNA sequences, MiRanda [40] and/or TargetScan [41] are often used. CircAtlas v2.0, circBase and circRNADisease also provide circRNA–miRNA interactions, without mentioning which miRNA database was used or how these interactions were predicted.

**CircRNA–RBP interactions**
In total, six circRNA databases (35%) report circRNA–RBP interactions. CircInteractome, CircRiC, CSCD and TSCD predict circRNA–RBP interactions based on crosslinking immunoprecipitation sequencing (CLIP-seq) data from starBase v2.0 [42]. The remaining two, CircFunBase and Circ2Disease, use the predicted RBP–circRNA interactions from CircInteractome (unfortunately offline).

**Protein-coding potential**
Although circRNAs are generally classified as noncoding RNAs, eukaryotic ribosomes can initiate translation of engineered circRNA when containing an internal ribosome entry site (IRES) element [26]. Furthermore, several human circRNAs are shown to be translated in vivo [43]. Therefore, some databases report predicted IRES or predicted open reading frames (ORFs). circRNADb provides the richest information on the protein-coding potential of circRNAs. It contains predicted IRES elements in the spliced sequence of each circRNA using VIPs (viral IRES prediction system) [44], and it also predicts the longest potential ORF. Other circRNA databases rely on CPAT (coding-potential assessment tool) [45] or ORF Finder (from NCBI), and IRESfinder [46] or IRESite [47] to predict the coding potential and IRES elements, respectively.

**CircRNA conservation**
As conservation of a particular genomic sequence may hint at a functional role, multiple researchers have investigated the conservation of circRNAs. CircAtlas v2.0, circbank and Circpedia v2 classify circRNAs from different species as orthologs when the BSJ site is conserved within a small 2–5 nucleotide range.

**Other annotations**
Finally, some circRNA databases have unique annotation features. For example, CircRiC reports the correlation between host gene expression and normalized BSJ read numbers. Furthermore, CircRiC also includes associations between drug response and the expression of circRNAs. MiOncoCirc v2.0 developed a pipeline (CODAC) to identify back-splicing involving two genes. Additionally, multiple databases report putative circRNA functions based on gene ontology enrichment analysis.

**Discussion**
There is little overlap among public circRNA databases
In total, 20 databases were included in this review, of which 14 noncurated and six curated. To assess the overlap among the circRNA databases, Euler plots were generated (Figure 2 and Supplemental Table 2).

The noncurated databases (Figure 2A) were divided into two groups, either based on de novo generated circRNA data or based on publicly available circRNA datasets (distinction also indicated in Table 1). First, for the databases that use publicly available circRNA datasets (circbank, circBase, circRNADb, CircInteractome, deepBase v2.0), we expect to see a high degree of overlap, as they often reuse the same datasets. While circbank, CircInteractome and deepBase v2.0 all use the circBase circRNA dataset as input, the contents of CircInteractome and deepBase v2.0 could not

| Database                | # circRNAs | Validation method** | Articles prior to | Short description                                                                 |
|-------------------------|------------|---------------------|-------------------|-----------------------------------------------------------------------------------|
| Circ2Disease [34]       | 249        | RT-(q)PCR microarray| 01 November 2017  | Reports curated circRNA-disease and circRNA-miRNA association                      |
| Circad [35]             | 925        | Unknown             | Unknown           | Reports curated circRNA-disease associations, including detailed experimental validation |
| CircFunBase [36]        | 3181       | RT-(q)PCR microarray RNA-seq | 01 May 2018 | Reports circRNA-associated miRNAs and circRNA-associated RBPs (CircInteractome) in seven plants and eight animals, including humans |
| Circ2Disease [37]       | 599        | RT-qPCR microarray  | 31 March 2018     | Reports curated circRNA-disease associations                                         |
| CircRNADisease [38]     | 328        | RT-qPCR microarray  | 01 November 2017  | Reports curated circRNA-disease associations                                         |
| LncRNADisease 2.0 [39]  | 736        | RT-(q)PCR microarray| 31 May 2018       | Reports curated circRNA-disease associations                                         |

*Number of unique human circRNAs in the download files.
**Most circRNAs in the database are validated by at least one of these methods. Some rarely used methods were omitted for clarity.
be assessed, as their databases were not online. Furthermore, although circbank contains more circRNAs than circBase, an observation we were unable to find an explanation for. circBase itself is based on nine circRNA datasets, including Jeck et al. [18] and Memczak et al. [19], two circRNA datasets that are also included in circRNADB (based on four datasets in total). Next, for the databases that use in-house or publicly available RNA-seq data (circAtlas v2.0, CIRCpedia v2, CircRiC, CSCD, exoRBase, MiOncoCirc v2.0 and TSCD), we observe little overlap. This is not unexpected as these databases rely on different samples, and circRNAs are expressed at low levels and with high sample-specificity [3,5]. Additionally, these circRNA databases applied different sequencing methods (varying RNA input levels, library preparation, sequencing dept, all affecting the sensitivity of the circRNA detection), circRNA detection tools and filtering steps, further contributing to the difference in content between the noncurated circRNA databases. The overlap between noncurated databases increases when filtered for experimentally validated circRNAs (circRNAs present in at least one curated database), as this also increases the probability of true positive circRNAs (Supplemental Figure 1).

It is thus extremely important to consider what samples were used to build the circRNA database and select a database in line with the tissue of interest. Moreover, the detection of circRNA in RNA-seq data does not guarantee that the predicted circRNAs are true positives. Therefore, some databases allow filtering for circRNAs detected by at least two tools, which improves the reliability of the predictions [48].

Overall, noncurated circRNA databases seem to contain a high number of circRNAs, whereby the reliability of these circRNAs must be questioned, as no validation using an orthogonal method is reported. Besides, it is important to recognize that circRNA expression and detection can vary considerably depending on multiple factors such as sample type, sequencing method and circRNA detection tool.

Six databases were found containing curated circRNA disease or function associations. Combined, these databases add up to 3522 circRNAs that have been empirically validated to date (Supplemental Table 3). Despite similar search strategies, there are notable differences in the content of curated databases (Figure 2B). Of note, not all databases apply the same criteria to label circRNAs as empirically validated. For the unpublished database circad, there is no information available on the accepted validation methods. Overall, the curated databases accept circRNAs validated by reverse transcriptase (quantitative) polymerase chain reaction (RT-(q)PCR), microarray or northern blot. However, CircFunBase and circRNADisease also accept RNA-seq as a sufficient method of circRNA validation. Seven hundred and forty four out of 3181 (23%) and 17 out of 328 (5%) circRNAs were solely detected by RNA-seq, respectively. It is not reported if a circRNA enrichment step (e.g. using RNAase R) was used as part of the RNA-seq validation, moreover both RNA-seq with and without circRNA enrichment can be found in...
the validated circRNAs when manually inspected. As a universal method for circRNA validation is lacking, we urge researchers to be more cautious and rely on multiple detection methods for effective validation of circRNAs.

Another explanation for the limited overlap between curated databases could be the redundancy arising from misannotated chromosomal positions. Usually, coordinates (chr:start-end) are given with inclusive start and end position. However, some formats (such as Browser Extensible Data, BED format) use a 0-based exclusive start position. Uncareful curation of circRNA positions from literature could thus result in incorrect or even redundant annotation. For example, there are two nearly identical CDR1 circRNAs present in CircFunBase, one with position chrX:139865340-139866824, and the other one with position chrX:139865339-139866824 (both hg19). The former is only supported by one publication, in contrast to the latter, which is supported by multiple publications and is also present in other databases (hsa_circ_0001946). However, these are probably the same molecules with BSJ positions based on different annotation systems.

In total, there are more than 2 million different circRNAs present in the union of all noncurated databases (compared to 384,066 predicted human RNA transcripts [49]), and 3522 circRNAs in the union of all curated databases. Surprisingly, more than 500 curated circRNAs are not present in any of the noncurated databases. This can partly be explained by misannotated start positions, as was previously discussed for CircFunBase. Although this issue does not completely explain these 500 circRNAs solely detected in curated databases, we expect that the remaining loss in overlap is due to other annotation related-issues. It is therefore recommended, when comparing datasets, to ensure that the annotations are compatible, or adjust them if necessary.

Further illustration of the lack in overlap can be seen in Supplemental Figure 2, which shows that most circRNAs are only present in one database.

The full-length sequence of circRNAs is lacking

Multiple databases (30%) provide full-length circRNA sequences and use it to predict interactions with miRNA and other sequence-based interaction partners. Unfortunately, the full-length sequence of most circRNAs is not known to date as full-length length circRNA sequencing datasets are lacking. Rather, the sequence between the start and stop position of the circRNA is inferred based on the reference genome sequence. The databases seem to report full-length circRNA sequences based on all known exons from the linear transcript. Most databases remove the introns, with the exception of circAtlas v2.0, which reports circRNA sequences based on exons and introns. This, however, relies on the unsupported assumption that circular and linear transcripts share the same splicing pattern and RNA sequence. However, almost 50% of the circRNA host genes give rise to multiple (up to 20) circular isoforms each [50]. Additionally, the inferred circRNA sequence depends on the genome build and on the exon and transcript annotation. CircFunBase reports multiple full-length circRNA sequences based on all overlapping linear transcripts reported by Ensembl, without taking into account alternative splicing.

In fact, all the databases included in this review should rather be called BSJ databases instead of circRNA databases, as none of them provide the empirically validated full-length sequence of circRNAs. An exception could be made for single-exon circRNAs, as it can be assumed that their sequence is the same as their parental linear exons. Interestingly, only 1.5% of circRNAs from the selected databases seem to be single-exon circRNAs. Whether this is caused by annotation problems in the different databases or is a true feature of circRNA biogenesis is unclear at this point.

While some tools, including CIRI-full [51] and circseq_cup [52], were developed to detect full-length circRNA sequences based on RNA-seq data, current databases do not make use of this type of analysis. CIRI-full makes use of a novel feature called...
reverse overlap and of the BSJ sites to reconstruct full-length circRNAs and circular isoforms. Circseq_cup first identifies BSJ sites and then assembles the full-length sequences of circRNAs using the paired-end reads aligned to the BSJ. Apart from computational methods, full-length circRNAs have been unambiguously identified using long-read single-molecule sequencing [53] or rolling circle amplification in combination with Sanger sequencing [54].

Another important characteristic of a circRNA is the DNA strand from which it is transcribed. Six of the databases we reviewed (30%) do not report the originating strand. Moreover, circRNADb does not report the gene nor the strand, but a link is provided to circBase, where the strand can be found. If the host gene is not mentioned, it is crucial to mention the strand from which the circRNA is transcribed (and hence stranded RNA sequencing methods should be used) to be able to identify the circRNA correctly.

### Ambiguous circRNA nomenclature contributes to the reproducibility crisis

Until now, no consensus circRNA nomenclature has been established. As indicated in Table 3, several similar nomenclature systems are in use, leading to multiple issues and increasing the risk of mistakes and confusion. The various nomenclature systems differ slightly in their prefix and the number of digits in the index. Next to the same circRNA having multiple names, nearly identical names with the same index sometimes correspond to different circRNAs. We illustrate this issue using circMTO1, which has at least 11 different names (Table 4). CircMTO1 (hsa_circ_0007874, hg19: chr6:74175931–74,176,329) is a circRNA that acts as a miRNA sponge for multiple RNA molecules, including oncogenic miR-9 [11] and linked to HCC [55]. circMTO1 is referred to as hsa_circ_30012 in circRNADb and hsa_circ_0007874 in circBase. It is problematic that hsa_circ_0030012 is a completely different circRNA in circBase, transcribed from FAM48A (hg19: chr13:37598171–37625720) and hsa_circ_07874 in circRNADb is also a completely different molecule (hg19: chr2:55040368-55047599). Similarly, while hsa-MTO1_00001 corresponds to a molecule with a BSJ at position chr6:74175932–74202075 in circAtlas v2.0, circcbank gives other BSJ coordinates to hsa_circMTO1_001, namely chr6:74175931-74176329 (matching the position mentioned in circBase). To further illustrate this issue, a second example presenting a list of all 13 names given to ciRS-7 (hg19: X_139865339:139866824) can be found in Supplemental Table 4.

Fortunately, most circRNA databases report the circRNA alias used in circBase, which was one of the first large circRNA databases. This is with the exception of circAtlas v2.0, circPedia v2, CircRiC, circRNADb, CSED and MiOncoCirc v2.0. The only way to compare the content of these databases is by using the BSJ position, which is not convenient.

Until now, the circBase nomenclature (hsa_circ_0000007) seems to be the most widely used naming system. However, it would be useful to work with a nomenclature that includes the host gene name, such as circcbank proposes. This makes the name more human-readable and can prevent mistakes. We recommend combining the species (e.g. hsa for human), the nature of the RNA molecule (circ), the official gene symbol (GENE), a then a unique identifier for each circRNA for that gene. While this unique identifier could be a simple three-digit number (from one to the total number of circRNAs identified...
for that gene), it would be more informative to have two indices (e.g. one to indicate the position of the BSJ and the second one to indicate the splicing pattern, once the full-length sequence of that specific circRNA is known). For example, two circRNAs from the same gene with the same BSJ, but with a different internal sequence, could be called hsa_circGENE_001_001 and hsa_circGENE_001_002. CircRNAs from which the full-length sequence is unknown, could be indicated by the _000 suffix. Alternatively, the strategy currently used by the miRNA database miRbase [56] can be applied, where instead of a second index a letter is used to indicate the full-length sequence. For example: hsa_circGENE_001a for the first known sequence and hsa_circGENE_001 for the unknown sequence. Using the gene symbol of the host gene poses another important issue: the naming system cannot be applied when the host gene lacks an official symbol, as is the case for many long noncoding RNAs (lncRNAs). Another unique identifier could be a hash that represents the full-length circRNA sequence or the 25 nucleotides flanking the BSJ if the full-length sequence is unknown. In any case, if a naming system without a host gene is used, it is crucial to report the strand from which the circRNA is transcribed, otherwise, the name can again refer to different circRNAs.

**CircRNA annotation is mostly based on assumptions and predictions**

Almost all circRNA databases report at least one type of circRNA annotation (*vide supra*), but these annotations should be handled with care. First, circRNA annotations are mostly based on computational predictions rather than experimental validation. Second, some sequence-based annotations, such as miRNA and RBP binding are predicted based on the presumed full-length sequence of circRNA molecules. As stated before, the empirically validated full-length sequence of a circRNA is generally lacking, and it is often not mentioned if the circRNA sequence used for prediction is the full-length sequence based on the reference genome, with or without taking splicing into account, or if the sequence is based on RNA-seq reads containing the BSJ. Third, it is important to note that the mere detection of a circRNA in a specific tissue or cell type does not necessarily indicate that this circRNA is specifically expressed. This also goes for curated circRNA databases, where the up- or downregulation of a circRNA in a specific disease sample in comparison with a control is often used to label a circRNA disease-specific.

Finally, a lot of databases do not report the source of their annotations and predictions. Overall, we would like to warn users of circRNA databases to be aware of the limitations regarding circRNA annotations.

**Updates, user interface and availability**

Although almost all authors of circRNA database articles mention the importance of regular updates and promise to maintain their database, only circBase and CIRCpedia v2 seem to have been updated after publication (in July 2017 and July 2018, respectively). Of note, some databases are very recent at the time of writing and might be updated in the near future. Unfortunately, some databases were completely inaccessible online, and some database exports were incomplete. Also, some databases are difficult to use or are limited to specific web browsers.

**Conclusions**

In this review, we provide an overview of all databases focused on human circRNA, divided into noncurated and curated circRNA databases. In total, there are more than 2 million different circRNAs present in the union of all noncurated databases, and 3522 circRNAs in the union of all curated databases. Generally speaking, there is limited overlap among these databases. The lack of overlap between noncurated databases can be explained by the use of different samples and the nature of circRNAs (low abundance, high sample-specificity) on the one hand, and by varying sequencing methods, circRNA detection tools and filtering on the other hand. It is important to be aware of the sample-specificity of circRNAs, and a database should be selected with care when conducting circRNA research. The lack in overlap among the curated databases might be due to different filtering techniques when selecting literature, and due to annotation-related issues. Furthermore, the use of different nomenclature systems is leading to redundancy and could cause confusion amongst circRNA researchers. This issue may very well contribute to the reproducibility crisis, and therefore we propose clear future guidelines for a solid circRNA nomenclature. Also, it is crucial to realize that the circRNA BSJ is not a unique identifier of a specific circRNA molecule, as splicing needs to be taken into account as well. Due to the lack of full-length circRNA sequences, multiple sequence-based annotations are predicted based on the assumption that circRNAs following the same splicing pattern...
as their parental mRNA counterparts and are thus unreliable. Finally, several databases are not regularly updated or suffer from connectivity issues.

**Key Points**

- There is limited overlap among circRNA databases, a result of the different source material and the nature of circRNAs (low abundance, high sample-specificity) on the one hand, and of varying sequencing methods, circRNA detection tools and filtering on the other hand.
- The ambiguous nomenclature of circRNAs resulted in conflicting names for circRNAs in different databases, contributing to the reproducibility crisis. One uniform naming system should be carefully implemented to prevent further future confusion.
- The BSJ position on itself is not sufficient to uniquely identify a circRNA, and the full-length sequence of most circRNAs is lacking.
- Many circRNA databases report interactions with miRNA and RBPs, protein-coding potential, conservation between species, etc. These predictions should be used with caution, as they are generally based on the assumed full-length sequence of circRNAs.

**Supplementary Data**

Supplementary data are available online at https://academic.oup.com/bib.

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