

TECHNICAL REPORT

CirInteractome: A web tool for exploring circular RNAs and their interacting proteins and microRNAs

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ABSTRACT

Circular RNAs (circRNAs) are widely expressed in animal cells, but their biogenesis and functions are poorly understood. CircRNAs have been shown to act as sponges for miRNAs and may also potentially sponge RNA-binding proteins (RBPs) and are thus predicted to function as robust posttranscriptional regulators of gene expression. The joint analysis of large-scale transcriptome data coupled with computational analyses represents a powerful approach to elucidate possible biological roles of ribonucleoprotein (RNP) complexes. Here, we present a new web tool, CirInteractome (circRNA interactome), for mapping RBP- and miRNA-binding sites on human circRNAs. CirInteractome searches public circRNA, miRNA, and RBP databases to provide bioinformatic analyses of binding sites on circRNAs and additionally analyzes miRNA and RBP sites on junction and junction-flanking sequences. CirInteractome also allows the user the ability to (1) identify potential circRNAs which can act as RBP sponges, (2) design junction-spanning primers for specific detection of circRNAs of interest, (3) design siRNAs for circRNA silencing, and (4) identify potential internal ribosomal entry sites (IRES). In sum, the web tool CirInteractome, freely accessible at <http://circinteractome.nia.nih.gov>, facilitates the analysis of circRNAs and circRNP biology.

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Introduction

Circular RNAs (circRNAs) are widely expressed RNAs lacking 5' and 3' ends, forming instead covalently closed RNA loops. In eukaryotes, circRNAs arise most often through backsplicing, a process in which the 5' and 3' ends of a spliced RNA are covalently linked to form a closed RNA molecule that contains exon and/or intron sequences.^{1–3} Although a few examples of circRNAs have been known for several decades, only with the recent widespread use of high-throughput RNA sequencing and bioinformatics have we learned that circRNAs constitute a vast class of stable RNAs expressed endogenously in cells, often with tissue-specific patterns.^{1–5} The functional roles of circRNAs are not as well established as those of other noncoding (nc)RNAs such as microRNAs (miRNAs). However, one prominent mechanism whereby circRNAs are believed to function is by sponging miRNAs, sequestering them away from protein-coding mRNAs.² It has also been postulated that circRNAs could serve as sponges for RNA-binding proteins (RBPs), platforms for assembly of RBPs, and protein-coding templates for translation.⁶

RBPs control all stages of post-transcriptional gene expression, including the splicing, export, turnover, translation, and localization of mRNAs.^{7–9} By modulating gene expression, RBPs play key roles in virtually all cellular processes – proliferation, differentiation, motility, senescence, apoptosis, as well as

the cellular responses to stresses, mitogens, and immune triggers.^{10,11} Consequently, RBPs have been implicated in a wide range of human diseases such as cancer, muscle pathologies, and neurodegenerative conditions.^{12–16} Recent developments in the analysis in RNA-protein (RNP) interactions using cross-linking techniques such as CLIP (cross-linking immunoprecipitation), PAR-CLIP (photoactivatable-ribonucleoside-enhanced CLIP), HITS-CLIP (high-throughput sequencing CLIP), and iCLIP (individual-nucleotide-resolution CLIP) have identified with unprecedented precision the binding sequences of RBPs on target RNAs.^{17–20}


MicroRNAs (miRNAs) comprise another major class of posttranscriptional regulators. They bind thousands of human transcripts through partial complementarity and generally reduce their expression by suppressing mRNA translation and/or stability.^{21–23} In some cases miRNAs may enhance mRNA translation.^{24,25} MicroRNAs are initially synthesized as primary (Pri)-microRNA transcripts, which are processed by the microprocessor complex into microRNA precursors (Pre)-microRNAs.^{26–29} Pre-microRNAs are then exported to the cytoplasm where they are further processed by DICER1 and loaded onto the RNA-inducible silencing complex (RISC) to be directed to target mRNAs.^{28–31}

RBPs and miRNAs can bind the same mRNA in a variety of functional manners; for example, they can compete, cooperate, or bind sequentially to a given mRNA.^{32–34} Such coregulatory effects by

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RBPs and miRNAs influence physiologic processes as well as disease.^{35,36} Thus, it is important to understand in detail their interaction with mRNAs. Several databases are available which facilitate research on RBPs (e.g., starBase) and miRNAs (e.g., TargetScan) interacting with specific mRNAs. Here, we have developed a new computational resource named ‘CircInteractome’ which enables researchers to search systematically for possible interactions of circRNAs with RBPs and miRNAs. This tool will greatly facilitate the identification of circRNAs that can sponge miRNAs and/or RBPs and will enhance the search for circRNAs with potential function as platforms for the assembly of RBPs. Additional features such as the ability to display the mRNA counterpart and the genomic and mature sequences of the circRNA, as well as the ability to design specific circRNA divergent primers and circRNA-directed siRNAs, are aimed at further facilitating other aspects of circRNA research. The CircInteractome tool currently searches against *Homo sapiens* databases but will be expanded to include other species in the future.

Results and discussion

CircRNA-wide mapping of RBPs and RBP ‘super-sponges’

Published CLIP datasets do not specify if the RBP binding sites are present in linear or circular RNAs (unless the CLIP hit spans a junctional sequence). Thus, we utilized CLIP datasets as indicated in the workflow (Fig. 1) to create a comprehensive binding map of RBPs to circRNAs (Fig. 2). We integrated 93 independently reported CLIP datasets from various RBPs (Table S1) obtained from different tissues and cell lines.³⁷ Computational analyses revealed that for select RBPs there were large numbers of binding sites in circRNA sequences (Table S2); for instance, we identified 117,000 circRNAs that could potentially associate with the RBP EIF4A3 (Fig. S1A). Analysis of other RBPs using CircInteractome indicated that they could also potentially interact with numerous circRNAs (Fig. S1A). For example, the mature circRNA hsa_circ_0000020 hosts multiple binding sites for several RBPs like HuR (6 sites) and FMRP (10 sites) (Fig. 2A, B). Thus, we hypothesized that circRNAs with relatively high density of binding sites for any single RBP could potentially act as a ‘sponge’ or a ‘decoy’ for that RBP. This sponging function would be enhanced by the long half-lives of circRNAs. By extension, circRNAs with exceptionally high density of binding sites for a given RBP might be considered to be ‘super-sponges’; for example, hsa_circ_0024707 (428 nt long) could function as a super-sponge for AGO2, since AGO2 can potentially bind this relatively short circRNA at 85 predicted positions (Table S2). In sum, this tool facilitates the search for potential RBPs interacting with circRNAs, and can identify possible RBP sponges as indicated in Figure 2 and Figure S3.

RBPs on circRNA junctions

RBPs may also interact with circRNA junctions and play a role in circRNA splicing, processing, folding, stabilization, and localization. To test this possibility, we queried CircInteractome for possibly binding sites for RBPs at select sequences spanning 100 nt upstream and downstream from the junction site (Fig. 2B). Computational analysis of the RBP binding sites from various datasets (Table S1) revealed that RBPs may indeed interact with circRNA junctions. For instance, EIF4A3 targets junction sequences with much higher frequency than the frequency seen for

targeting the body of mature circRNAs (Fig. S1B). This suggests that EIF4A3 could have a preference for binding to circRNA junctions compared to other RBPs (Fig. S1B). An example of this type of search for hsa_circ_0000020 is shown in Figure 2B and Figure S4A.

Mapping binding sites of RBPs on pre-circRNA

The splicing machinery can generate circRNAs through non-linear back-splicing, which joins 5' and 3' ends covalently to make a circRNA.³⁸ Splicing events are tightly regulated by RBPs and snRNAs binding near the splice sites.³⁹ Thus, we used CircInteractome to search all datasets (Table S1) in order to identify the binding sites of RBPs in the flanking sequences upstream and downstream of the mature circRNA. For instance, the flanking sequences on both sides of the hsa_circ_0000020 junction indicated 4 binding sites for EIF4A3 (Fig. 2C, bottom; Fig. S4B), suggesting a possible role for EIF4A3 in the biogenesis of hsa_circ_0000020. Our analysis also uncovered a relatively higher frequency of putative binding sites for splicing factors like TDP43 at the flanking sequences of circRNAs, which point to a possible role for these RBPs in circRNA splicing/biogenesis (Fig. S1C).

Potential circRNA translation though IRES

CircRNAs are believed to not be translated. While most circRNAs do not appear to be associated with polyribosomes, the possibility exists that some circRNAs might be translated into protein products.³ In this regard, linear long noncoding (lnc) RNAs are not generally translated into proteins, but a subset of them appear to be a source of functional small or micropeptides due to the presence of short open reading frames (ORFs).⁴⁰ For example, a micropeptide encoded by a noncoding RNA was recently found to regulate muscle performance.⁴¹ In addition, the viral circRNA CCC (covalently closed circular, 220 nt long) was recently found to be fully translated into a 16-kDa highly basic protein in infected rice plants.⁴² Bioinformatic analysis revealed the presence of IRES (internal ribosome entry sites) and sites for RBPs that regulate IRES-mediated translation (IRES trans-acting factors or ITAFs) in several circRNA sequences (Table S3).⁴³ For example, we found that hsa_circ_0041407 contains an IRES (underlined sequence) and partial coding sequences of MAX network transcriptional repressor (*MNT*; Fig. 3A, B) and postulated that hsa_circ_0041407 could give rise to a small chimeric protein of ~31 kDa (highlighted sequence, Fig. 3B). To our surprise, IRES regions of circRNAs are predicted sites for many RBPs, including HuR and PTB, 2 proteins reported to modulate IRES-driven translation.⁴³ Together, these findings suggest that circRNAs could be translated through IRES sequences, and thus IRES-bearing circRNAs detected in association with polyosomes warrant future exploration.

miRNA ‘super-sponge’ circRNAs

The main function described for circRNAs thus far in the literature is that of miRNA sponges, as shown for the circRNA ciRS-7, which has multiple binding sites for miR-7.⁵ To

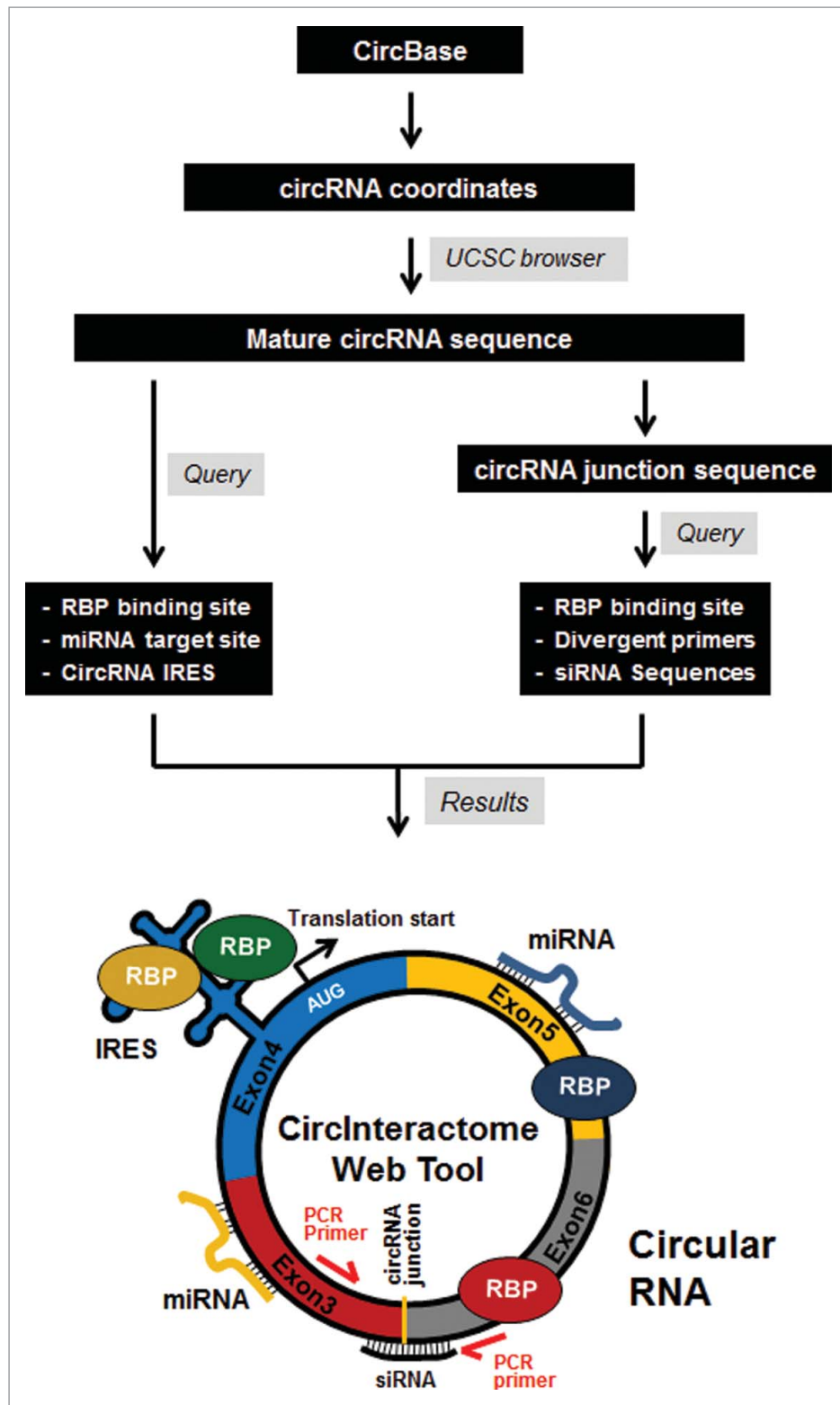


Figure 1. Workflow of the web tool Circular RNA Interactome or 'CirInteractome'.

characterize miRNA-circRNA interactions (Fig. 4, Fig. S5), we incorporated into CirInteractome the ability to search using the TargetScan algorithm, which predicts miRNAs that target circRNA by surveying for 7-mer or 8-mer complementarity to the seed region as well as the 3' end of each miRNA.⁴⁶ A survey of miRNA target sites in circRNAs revealed the presence of

numerous target sites for a specific miRNA. In cases of exceptionally high hit numbers, a circRNA could be considered a 'super-sponge' for microRNAs. Strikingly, over 3,000 circRNAs were found to have at least 20 miRNA target sites in a single circRNA, and most of them had AGO2 binding sites (Table S4). For example, hsa_circ_0139850, which is only

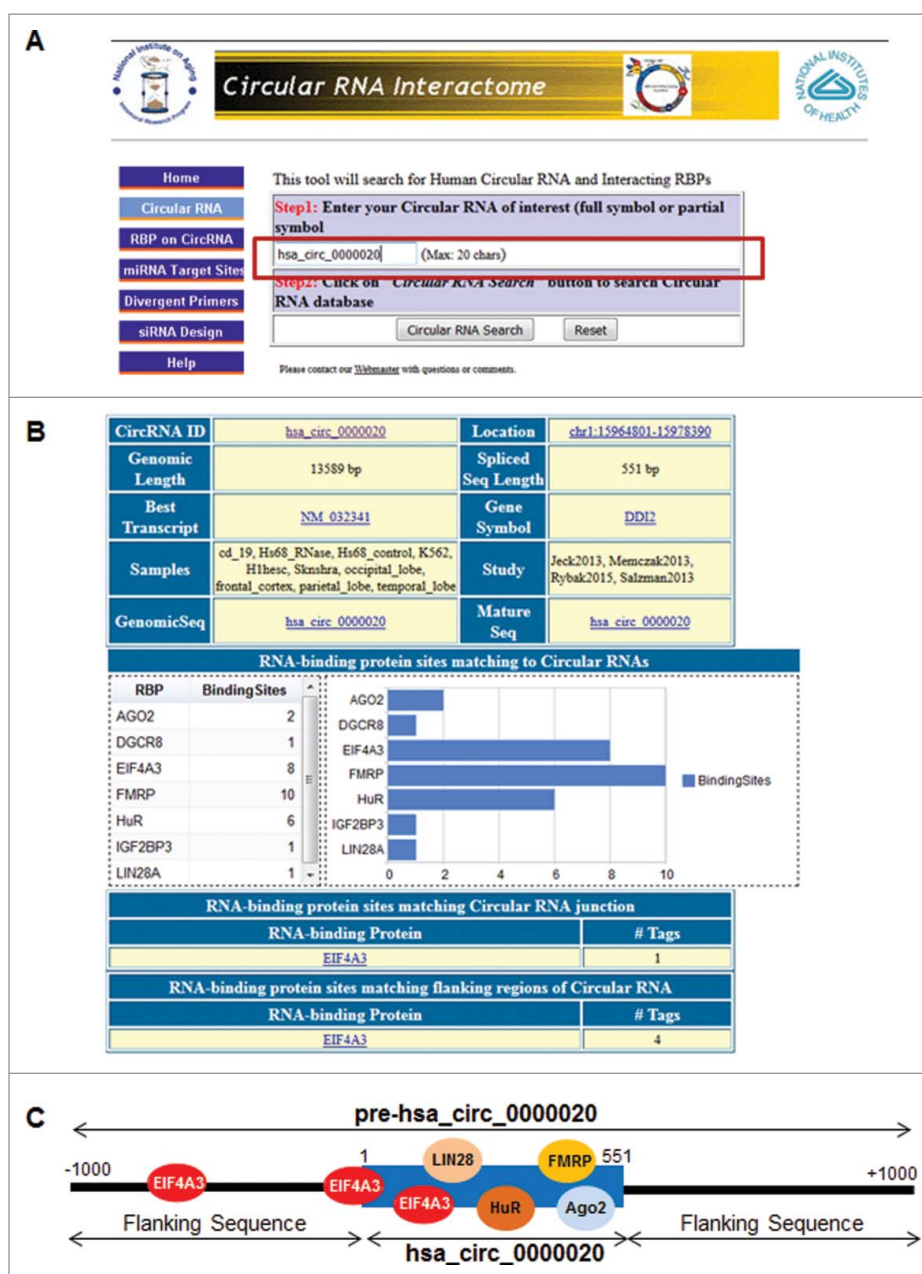


Figure 2. View of CircInteractome input and output pages. **A.** Example the 'Circular RNA' page exhibiting the input parameters needed for a CircInteractome run. **B.** Screenshot of 'Circular RNA search' for hsa_circ_0000020, showing RBPs binding in different regions of this circRNA. **C.** Schematic representation of the potential binding sites of different RBPs to mature and pre-hsa_circ_0000020.

437 nt long, has 22 sites for miR-7 and thus might act a super-sponge for miR-7 (Fig. 4). The general finding that circRNAs often have more miRNA target sites than those predicted by chance lends further support to the idea that binding microRNAs could be one of the key functions of circRNAs.

Divergent primer design

Designing specific primers for quantification of circRNA using qPCR amplification can be challenging and prone to errors, since the mature circRNA sequences after splicing are not readily available in many cases and the primers must be divergent and must span the junction. At present, there is no software or web server that enables direct design of divergent primers

specific to circRNAs. Thus, we incorporated primer design tools into CircInteractome (Primer3⁴⁵ or NCBI primer design tool) and used as template the sequence around the circRNA junction to ensure that the circRNA was amplified specifically by reverse transcription (RT) followed by real-time quantitative (q)PCR analysis (Fig. 5A). To illustrate this feature of CircInteractome, we show the design of divergent primers at the junction sequence of hsa_circ_0000020 (Fig. 5B; Fig. S6A,B).

siRNA design

Designing siRNAs selectively directed at circRNAs is challenging due to the limited knowledge of the junctional sequences and the lack of appropriate software

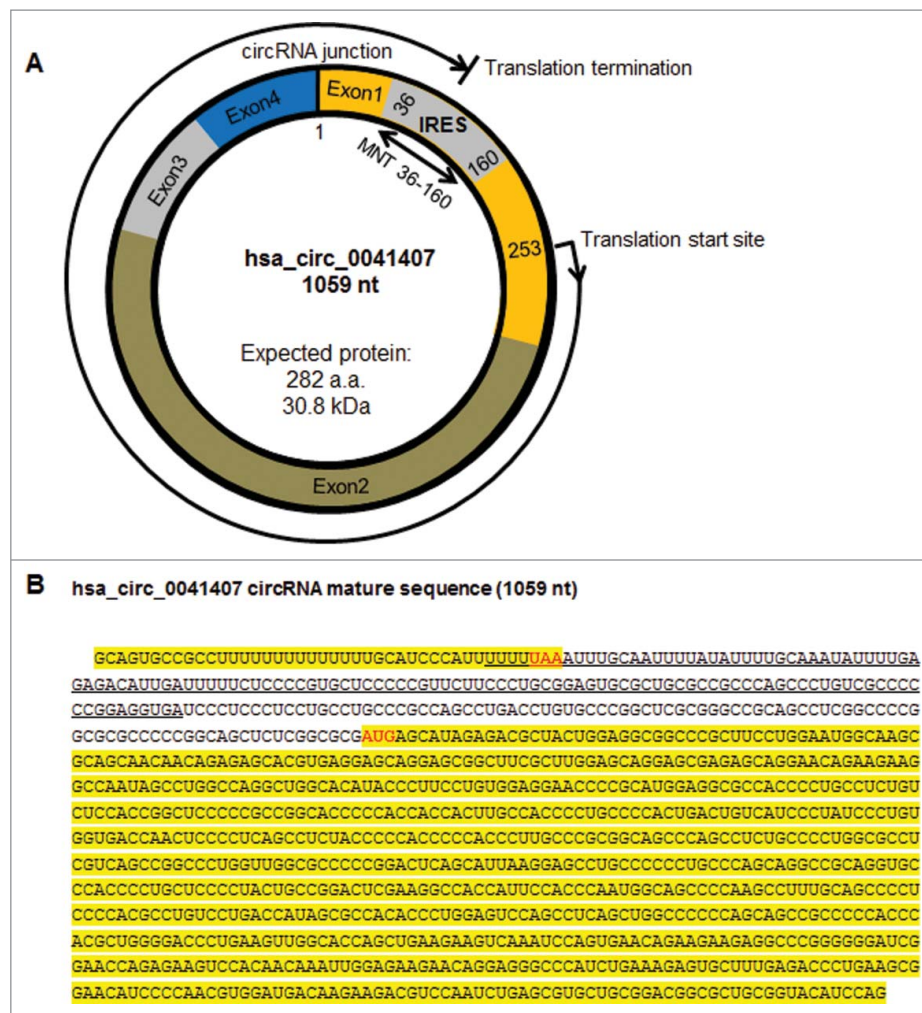


Figure 3. Predicted IRES possibly mediating circRNA translation. A. Schematic representation of hsa_circ_0041407, which contains the *MNT* 36-160 IRES that may potentially drive translation. B. The sequence of hsa_circ_0041407 highlighted can potentially be translated via the IRES underlined.

tools to help design them. Thus, we incorporated into CircInteractome the ability to generate 21-nt siRNAs targeting junctional sequences spanning 5–16 nt on either side of the junction. Using standard siRNA design criteria, we filtered junctional sequences to find the best siRNA target sequence^{46,47}; users need to avoid siRNAs with long stretches rich in G/C, and confirm using BLAST that the siRNAs are specific. For better silencing effects, the output siRNAs may be ordered with an additional 2 nucleotides (dTdT) as 3' DNA overhangs (Fig. S7). Users may then purchase them from any source; links to forms to complete siRNA orders using IDT or Dharmacon are provided. Examples of the 10 best possible siRNA target sequences for hsa_circ_0000020 are shown (Fig. 6). In short, CircInteractome can be used to predict siRNA sequences targeting the circRNA junctions.

Conclusions and future directions

This freely available web service can facilitate the study of circular RNAs and their interactions with other binding factors, mainly RBPs and miRNAs. CircInteractome provides researchers with valuable detail about circular RNAs and their possible

role in sequestering RBPs and/or miRNAs and thereby reducing their availability for mRNAs. CircInteractome also facilitates the design of primers for studying circRNA by RT-qPCR analysis. CircInteractome can be used to predict RBP binding to upstream and downstream sequences of the pre-spliced transcript, thus potentially shedding light into the biogenesis of circRNAs. The interaction of translation regulators (miRNAs, RBPs, ITAFs) and the presence of IRESs in circRNAs can help to decipher whether circRNAs have protein/peptide-coding capabilities.

CircInteractome incorporates several features from other freely available web resources, such as circBase, StareBase 2.0, TargetScan 7.0, and Primer3. By integrating these resources, CircInteractome enables the user to find out the genomic and mature circRNA sequences (Fig. S2), circRNA-binding partners (RBPs, miRNAs), siRNAs, and primers to study circRNA levels, localization, and function.

Although CircInteractome has many user-friendly features for circRNA researchers, it is limited in its ability to predict RBP and miRNA interactions when circRNAs form secondary or tertiary structures. Given that all of the data provided here are predicted based on sequence matches and the presence of

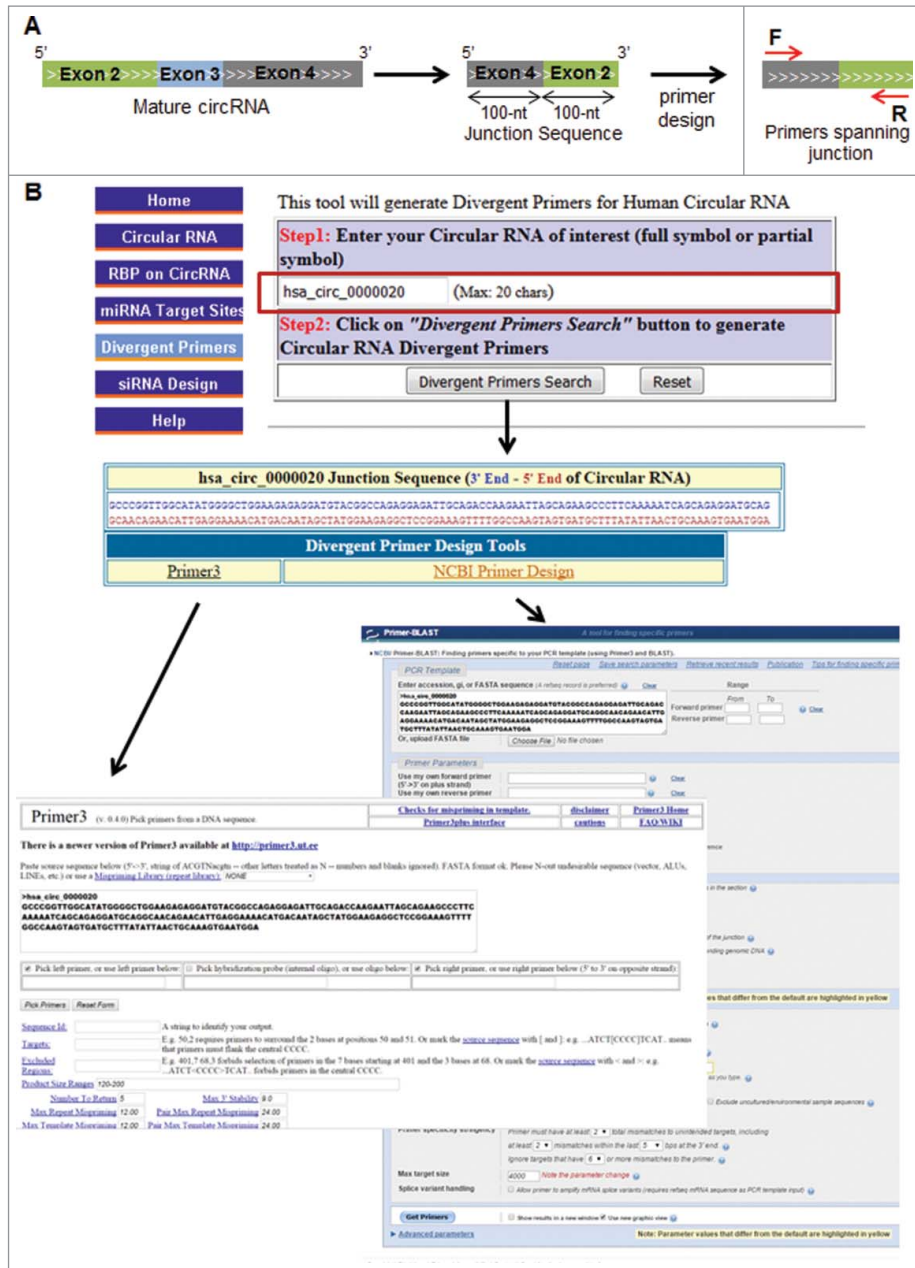


Figure 5. CircRNA primer design. A. Schematic representation of the design of divergent primers for circRNA detection by RT-qPCR analysis. B. Input webpage for divergent primer design. C. Screenshot of the output page showing the junction sequence of circRNA and links of primer design tools (Primer3 and NCBI).

design (Fig. 5A). This dataset was used for sequence alignment with RBP binding sites.

Acquisition of binding sites of RBPs on circRNAs

Datasets including the binding sites of 35 RBPs (Table S1) identified by PAR-CLIP, HITS-CLIP, or iCLIP were retrieved from starBase v2.0 (<http://starbase.sysu.edu.cn/>),³⁷ UCSC browser, and published datasets. A complete list of all datasets used in the study is shown in Table S2. Data were previously analyzed and thus we used these datasets for our analysis without modification. The sequences for the RBP CLIP clusters were downloaded from the UCSC browser mirror (<http://genome.mdc-berlin.de>)⁴⁹ and were analyzed against circRNA mature, flanking, and junction sequences of circRNAs with 95% sequence

homology and maximum of 2 mismatches to find their binding sites (Fig. 1). When more than one tag was predicted at the same site for an RBP, only one tag was considered as the binding sequence.

IRES dataset

The sequences of reported IRES were downloaded from the IRESite (<http://iresite.org/>), which contains information about experimentally validated IRES sequences.⁵⁰ These IRES sequences were analyzed against mature circRNA sequences to find out the presence of IRES in circRNAs; 99% similarity between the IRES sequence and the mature circRNA sequence was allowed (Fig. 1). IRES sequences present in circRNAs were further analyzed against the RBP CLIP cluster sequences to

Figure 6. CircRNA siRNA design. Screenshot of the CirInteractome siRNA design page, including an example of output siRNAs targeting the junction sequence of a given circRNA.

identify binding sites of different RBPs in the IRES sequences of circRNAs (Fig. 1).

miRNA Target circRNAs

Mature sequences of circRNAs were used in the TargetScan Perl Script to predict the miRNAs which have sequence complementarity with circRNA.⁴⁴ The complete miRNA list and sequences were taken from the microRNA database (<http://www.mirbase.org/>).⁵¹

Divergent primer design

The circRNA junction sequences were retrieved from the mature circRNA sequences and exported to primer design tools (Primer3 or NCBI) to give maximum of 5 sets of primer pairs for PCR products of sizes ranging from 120–200 bp and spanning the junction (Fig. S6A, B). For circRNAs smaller than 200 nt, users are given a choice for the length of PCR amplicon based on circRNA length.

CircRNA siRNA design

siRNAs spanning circRNA junction were designed using criteria for 21-nt siRNAs described previously.^{46,47} To design the 21-nt siRNA target sequence spanning the junction, we used a minimum of 5 nt on either side of the junction.

Availability

The database is freely accessible through the web server at <http://circinteractome.nia.nih.gov>.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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