Integrative Epigenomic Analysis of Transcriptional Regulation of Human CircRNAs

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Circular RNAs (circRNAs) are evolutionarily conserved and abundant non-coding RNAs whose functions and regulatory mechanisms remain largely unknown. Here, we identify and characterize an epigenomically distinct group of circRNAs (TAH-circRNAs), which are transcribed to a higher level than their host genes. By integrative analysis of cistromic and transcriptomic data, we find that compared with other circRNAs, TAH-circRNAs are expressed more abundantly and have more transcription factors (TFs) binding sites and lower DNA methylation levels. Concordantly, TAH-circRNAs are enriched in open and active chromatin regions. Importantly, ChIA-PET results showed that 23–52% of transcription start sites (TSSs) of TAH-circRNAs have direct interactions with cis-regulatory regions, strongly suggesting their independent transcriptional regulation from host genes. In addition, we characterize molecular features of super-enhancer-driven circRNAs in cancer biology. Together, this study comprehensively analyzes epigenomic characteristics of circRNAs and identifies a distinct group of TAH-circRNAs that are independently transcribed via enhancers and super-enhancers by TFs. These findings substantially advance our understanding of the regulatory mechanism of circRNAs and may have important implications for future investigations of this class of non-coding RNAs.

Keywords: transcription factors, transcriptional regulation, epigenomic, circular RNAs, TAH-circRNAs

INTRODUCTION

CircRNAs are a large class of non-coding RNAs produced by circularization of exons, and they are characterized by the presence of a covalent direct ligation of 3' and 5' ends generated by back splicing (Chen, 2016). Although the vast majority of circRNAs have not been studied, several lines of evidence have suggested that this group of non-coding RNAs play important roles in tissue development (You et al., 2015), gene regulation, and carcinogenesis (Hsiao et al., 2017). With the advent of high-throughput profiling technologies, a myriad of circRNAs have recently been identified across various types of cells, tissues, and species (Zhang X. O. et al., 2016; Dong et al., 2017). Intriguingly, although the expression levels of the majority of circRNAs are expectedly low,
some of them are transcribed at comparable or even higher levels than their linear counterparts (Li X. et al., 2017). For example, circRNAs are highly abundant in the fly brain (Westholm et al., 2014), mammalian neuronal and muscle tissues (Rybak-Wolf et al., 2015).

Although it is generally assumed that circRNAs are the spliced products of their host transcripts, a few lines of evidence suggested that some circRNAs may be transcriptionally activated independently from their host genes. For example, a previous study found that expression changes of a group of circRNAs were largely independent of host genes during the differentiation of epidermal stem cells (Kristensen et al., 2018). Moreover, the transcription factor Twist1 selectively promoted the transcription of circRNA Cul2 but not its host gene (Meng et al., 2018). In addition, Bin et al. observed that circRNA Nfix was driven by a super-enhancer (SE) in cardiomyocyte cells, which was also independent from its host gene (Huang et al., 2019). Interestingly, during neuronal differentiation, some circRNAs were up-regulated to a higher extent compared to their host mRNAs (You et al., 2015). However, it is unclear whether these observations represent merely particular cases or there exists a general regulatory mechanism for independent transcription of circRNAs.

To address this question, we performed comprehensive epigenomic analyses of circRNAs and identified a distinct group of these non-coding RNAs that are transcriptionally activated to a higher level than the host genes (TAH-circRNAs). Specific TFs occupy genomic regions of these TAH-circRNAs, which exhibit higher conservation than other circRNAs. Cis-regulatory regions of TAH-circRNAs have increased active histone modifications and reduced DNA methylation. Furthermore, TAH-circRNAs associated with SE have higher expression level and denser TF occupancy than other circRNAs. Lastly, we functionally validated a group of TAH-circRNAs that were independently transcribed by a master regulator TF, FOXA1. These results together provide many new insights into transcription regulatory mechanism of circRNAs.

**MATERIALS AND METHODS**

**Cell Culture**

HepG2 cells were maintained in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum (FBS). Cells were incubated at 37°C in a 5% CO₂.

**siRNA Transfection**

Three different siRNAs against human FOXA1 (Supplementary Table 3) were designed and ordered from GenePharma (Shanghai, China). Synthetic sequence-scrambled siRNA was used as a negative control (NC). HepG2 cells were seeded in six-well plate until cell density reached 30–40% prior to siRNA transfection. Three siRNAs were mixed and transfected using Lipofectamine RNAiMAX (Invitrogen, USA) and incubated for 48 h.

**Real-Time PCR Detection**

Total RNA was isolated from HepG2 cells using TRizol (Invitrogen, USA) and was reverse-transcribed to cDNA with RevertAid RT Reverse Transcription Kit (Thermo Scientific, USA). Real-time PCR was performed with SYBR Green I Master (Roche, USA) in LightCycler 480 System. The sequences of the PCR primers were listed in Supplementary Table 3. All samples were measured in triplicate and normalized by GAPDH. The data were expressed as the mean ± SD of three independent experiments.

To validate the sequences of circRNAs in HepG2 cells, the products of real-time PCR were sequenced by Sanger method and the results completely matched with the predicted sequences.

**Microarray**

Agilent Human V6 (4*180K, Design ID: 084410) was used for hybridization of total RNAs and was then scanned by Agilent Scanner G2505C (Agilent Technologies, USA) to measure mRNA and circRNA.

**Luciferase Reporter Assay**

Hsa_circ_0061137_upstream + 1,500 nt and hsa_circ_0061137_upstream + 1,000 nt were cloned into pScheck2 luciferase reporter vector (Promega). Luciferase vector and control vector were transfected into HepG2 cells with Lipofectamine 3000 (Thermo Fisher Scientific). After 48 h of transfection, the reporter activity was measured by the Dual-Luciferase Reporter Assay System (Promega, E1910).

**Genomic Annotation of CircRNAs and Coding Genes**

Genomic annotations of 92,375 human circRNAs and 20,287 coding genes were downloaded from circBase database (December 15, 2015, http://www.circbase.org/) and ensemble (GRCh37, http://grch37.ensembl.org/index.html), respectively.

**Identification of CircRNAs and Quantification of the Expression Levels of CircRNAs and Coding Genes**

We downloaded ENCODE RNA-seq data of 36 cell lines, including PolyA- RNA-seq data of 17 cell lines and Total RNA-seq data of 19 cell lines (Supplementary Table 1). The find_circ (Memczak et al., 2013) computational pipeline was applied to identify circRNAs for each cell line. These identified circRNAs were next annotated to circBase database. A circRNA was assigned to the corresponding circBase ID if it had the same positions with an annotated circRNA in the database. We computed the expression level of circRNAs by different methods: (1) CircRNAs expression was normalized as the number of backsplice junction spanning reads per million raw reads (RPKM). Since longer reads will have more power in detecting backsplicing events, read length can be used as normalization for circRNAs. Because find_circ software requires anchor sequences of 20 bp on each side of the read, a reads per kilobase per million mapped reads (RPKM) expression value for each circRNA was calculated by dividing circRNAs RPM value by 112 and multiplying by 1,000 (Veno et al., 2015).
(2) We computed the expression of gene body of circRNAs using the following equation:

\[
\text{RPKM}_{\text{body}} = \frac{10^9 \times m}{n \times L}
\]

Where \(m\) corresponds to the number of reads locating in gene body of circRNAs; \(n\) corresponds to the total number of reads; and \(L\) represents the length of circRNAs.

(3) H3K27ac modification is associated with the regulatory regions of active genes. Therefore, we considered that if the proximal regulatory region of a circRNA has higher H3K27ac signals than other circRNAs, it is likely to have higher transcriptional activity. To measure the transcriptional activity of circRNAs, we normalized H3K27ac signals within proximal regulatory region of circRNAs, which is defined as a region that spread 1,000 bp upstream and 1,000 bp downstream from its TSS. The transcriptional activity of a circRNA was measured as:

\[
\text{RPKM}_{\text{ss}} = \frac{10^9 \times C}{2000 \times N}
\]

Where \(C\) corresponds to the number of H3K27ac reads locating in regulatory region of the circRNA; \(N\) corresponds to the total number of reads.

**Genome-Wide Binding Analysis of TFs**

A total of 690 TF ChIP-seq data were downloaded from ENCODE. The peak list of these ChIP-seq data was based on 161 unique TFs, generated from uniformly processed pipeline. This dataset spans 91 human cell types, where the calling method of SPP (Kharichenko et al., 2008) was used to identify the regions of enrichment (peaks) ([Supplementary Table 1](#)). In addition, we used ChIP-seq data of histone marks and DNaseI.

Moreover, we analyzed ChIP-seq data of H3K27ac histone mark from Hnisz et al. (2013), which included eight (GM12878, H1-hESC, HeLa-S3, HepG2, HUVEC, IMR90, K562, MCF-7) cell lines.

**DNA Methylation Analysis**

We downloaded processed beta-values of DNA methylation 450 k array (61 cell lines) and whole-genome bisulfite sequencing (WGBS) data (four cell lines) from ENCODE. We converted the genome coordinate of WGBS (GRCh38) data to hg19 through liftover (https://genome.ucsc.edu/cgi-bin/hgliftOver). For WGBS dataset, only CpGs with a minimal coverage of 4 were included for analysis.

**Construction of TF-Mediated Regulatory Network of CircRNAs and Coding Genes**

For each of 92,375 circRNAs from circBase, we identified those TFs that occupy the regulatory regions of circRNAs through searching 161 TF ChIP-seq data from ENCODE. A circRNA is considered as occupied by a TF if its regulatory region (−2/+1 Kb of its TSS) has at least one peak. We merged all the identified TF-circRNA pairs to a global regulatory network of TF-circRNA. In parallel, we also constructed regulatory network of TF-coding genes by the method. According to the global regulatory network of TF-circRNA, we abstracted a regulatory sub network of TAH-circRNAs through mapping TAH-circRNA and abstracting nearest TFs of them.

**ChIP-Seq Data Processing and Identifying ChIP-Seq Enriched Region in Human Cells**

Sequencing reads were aligned to human genome build hg19 using bowtie 0.12.9 using parameters -K 2 -m 2 -n 2 -s. Enrichment regions of H3K27ac and TF in human sample were calculated using MACS 1.4.2 (Zhang et al., 2008; Jiang et al., 2019; Qian et al., 2019; Tang et al., 2019; Feng et al., 2020; Li et al., 2020) using parameters --keep-dup 1 --wig --single-profile--space = 50 -p 1e-9 on H3K27ac and various TF ChIP-seq with control libraries. UCSC Genome Browser tracks were generated with MACS wiggle outputs. MACS peaks of human H3K27ac were used as constituent enhancers for SE identification. ChIP signal visualization was done by IGV (Integrative Genomics Viewer) (Thorvaldsdottir et al., 2013) and the input WIG files were created by MACS.

**Identification of SE and Associated CircRNAs**

SE were identified using ROSE (https://bitbucket.org/young_computation/rose) with parameters -s 12,500 -t 2,000 by stitching enhancers, which has been described previously (Loven et al., 2013). Typical-enhancer and SE associated circRNAs were identified using a Python script of ROSE.

**CircRNA Microarray of HCC Samples**

We re-annotated and re-analyzed the circRNA expression profiles of HCC sample from GEO GSE97332, which was based on Agilent circRNA microarray. Specifically, we created a library of circRNA junction sequences containing 45 bp from tail and 45 bp from head of each circRNA in circBase. We then performed re-annotation by aligning the probe sequences of the Agilent circRNA microarray to our customized junction library with BLASTn tools (McGinnis and Madden, 2004). Results from the sequence alignment were further processed as follows: (1) we retained the probes matching to only one unique circRNA, resulting in a set of probe-circRNA pairs. (2) Each circRNA is required to be perfectly matched to at least one probe. As a result, a total of 5,515 circBase circRNAs were re-annotated.

**ChIA-PET**

We downloaded the RAD21, CTCF, POLR2A, and ESR1 ChIA-PET data in three cell lines (HepG2, K562, and MCF7) from ENCODE. We then analyzed these datasets by ChIA-PET2 (Li G. et al., 2017) using parameters -e 1 -k 2. In addition, we downloaded the chromatin interaction data of GM12878 and IMR90 from 4DGenome and extracted the chromatin interactions that were detected by 3C, 5C, and Hi-C.

**Gene Set Enrichment Analysis (GSEA)**

GSEA was performed using the tool from http://software.broadinstitute.org/gsea/index.jsp. In brief, \(t\)-test in gene expression from two experimental conditions was calculated. The result was used as a ranked list in the Pre-Ranked function.
Identification of a Group of TAH-CircRNAs With Distinct Epigenomic Features

To comprehensively determine the expression profiles of circRNAs, we first annotated all circRNAs in 36 cell lines (with matched Total RNA-seq and Poly-A RNA-seq data) using the Find_circ software, and computed the RPKM (Reads Per Kilobase Million) values of both circRNAs and the host genes. We then focused on six cell lines (GM12878, H1-hESC, K562, HepG2, IMR90, and MCF7) from ENCODE consortium since they have comprehensive epigenomic data, including profilings of TF occupancies, histone modifications, and DNA methylation. We found that a significant proportion of circRNAs (40–60% across different cell lines) had higher RPKM compared to their host genes, in line with previous reports (Salzman et al., 2013; Rybak-Wolf et al., 2015; You et al., 2015; Zhang Y. et al., 2016; Legnini et al., 2017; Kristensen et al., 2018). Moreover, this increase over host genes was only observed in circRNAs specifically identified by Find_circ but not in predicted circRNAs using circBase, suggesting the specificity of the expression difference between circRNAs and host genes. As examples shown in Figure 1C, the level of hsa_circ_0001727 was ∼15 times higher than its host genes, respectively. Indeed, as much as 6–24% of circRNAs had at least 6-fold higher expression levels than their hosts (Figure 1D). At the same time, we calculated the number of TAH-circRNAs exons and their host genes in six cell lines, respectively (Supplementary Figure 7). The distribution of TAH-circRNAs exons was different from their host genes. Most of TAH-circRNAs contained one or two exons, whereas their host genes were composed of multi-exons. For instance,
hsa_circ_0001727 was originated from exon 2 and exon 3 of the host gene (contained six exons, ZKSCAN1), and the expression level of this TAH-circRNA was higher than its host gene, consistent with the result found by Yao et al. (2017). Hsa_circ_0001900 originated from exon 2 and exon 3 of the host gene (contained 17 exons, CAMSAP1). Zhu et al. measured circ_0001900 and found that the circRNA was significantly up-regulated in CRC tissues and much more stable than the host gene (CAMSAP1) (Zhou et al., 2020). Taken together, these results indicated that TAH-circRNAs exhibit the different transcription pattern of exons compared with host gene.

To explore the transcriptional activity of circRNAs with higher expression levels than the hosts, we next analyzed H3K27ac (a histone marker indicative of active transcription) ChIP-seq profiles from these six cell lines. Specifically, we measured H3K27ac intensity in potential proximal regulatory regions of circRNAs (between 2 Kb upstream and 1 Kb downstream of the circRNA TSS). Consistent with the changes in RNA expression, we observed that 30–40% circRNAs had higher H3K27ac signals than the host genes (Supplementary Figure 1H). Importantly, circRNAs with higher H3K27ac signals strongly overlapped with those having higher expression levels (Figure 2A, Supplementary Table 2). In fact, 17–32% of circRNAs with higher H3K27ac signals had higher expression levels, strongly suggesting that the increased level of abundance of these circRNAs is not simply due to back-splicing. We considered that circRNAs with both higher H3K27ac signals and elevated expression levels might potentially have unique transcriptional mechanism, and thus we focused on characterizing this group, which we termed TAH-circRNAs (Transcriptionally Activated to a Higher level than host genes). TAH-circRNAs accounted for 20–30% of all circRNAs across various cell lines (Supplementary Figure 2A). Importantly, a significantly positive correlation between the expression level and the H3K27ac intensity was observed only in TAH-circRNAs (Figure 2B), but not in non-TAH-circRNAs. Furthermore, TAH-circRNAs had both higher H3K36me3 signals and POLR2A loadings than other circRNAs in each cell line (Figure 2C, Supplementary Figures 2B,D–F). Consistently, TAH-circRNAs displayed more elevated DNaseI signals than non-TAH-circRNAs (Supplementary Figure 2C). Lastly, we observed that TAH-circRNAs were more evolutionarily conserved than other circRNAs (Figure 2D), implying that TAH-circRNAs might have more important functions and thus were more conserved across species. Figure 2E shows one example of TAH-circRNAs (hsa_circ_0061137), which had higher levels of H3K4me1, H3K36me3, as well RPKM values than the host gene (DIDO1). Most importantly, ChiA-PET results identified direct interactions between the TSS of this circRNA and two distal cis-regulatory regions. These two distal regions had conspicuous H3K27ac modification, DNaseI signals, as well as POLR2A binding, indicative of enhancer function. Furthermore, FOXA1 and CEBPB were found to occupy both its TSS and one of the cis-regulatory regions, implying that this circRNA might be under direct transcriptional regulation from these two TFs. Other example TAH-circRNAs are shown in Supplementary Figures 6A–D. The existence of hsa_circ_0061137 promoter was further experimental verified by luciferase reporter assay in HepG2, which showed the high transcriptional activity of hsa_circ_0061137 transcriptional start site (upstream 1,500/1,000 nt) (Figure 2F). Taken together, these analyses distinguished TAH-circRNAs from other circRNAs in terms of epigenomic and biological features.

### Regulatory Regions of TAH-CircRNA Have Low DNA Methylation

DNA methylation plays a fundamental role in regulating gene expression, and it is well-established that an inverse relationship exists between DNA methylation and transcriptional activity at cis-regulatory regions, such as enhancers and promoters (Lister et al., 2009; Silva et al., 2019). CircRNAs with hypo-methylated regulatory regions were occupied by more TFs than those with hyper-methylated regions across multiple cell lines, indicative of more active TF regulation (Figures 3A,B and Supplementary Figures 3A,B). Notably, DNA methylation was significantly lower in regulatory regions of TAH-circRNAs relative to other circRNAs (Figure 3C and Supplementary Figure 3C), suggesting that the hypo-methylated regulatory regions in TAH-circRNAs are more accessible for TF binding and regulation. This methylation pattern is consistent with our earlier characterizations of other epigenomic features of TAH-circRNAs, displaying more accessible regulatory regions, stronger transcriptional activities, as well as higher expression levels.

### Comprehensive Analysis of TF-CircRNA Regulatory Network

To further understand the transcriptional regulatory mechanisms of TAH-circRNA, we identified TFs that had binding sites within the potential proximal regulatory region of TAH-circRNAs (between 2 Kb upstream and 1 Kb downstream of the TSS) by analyzing 159 TF ChIP-seq results from the matched six cell lines (ENCODE database). We first analyzed the distribution degree of TF-circRNAs regulatory network, and determined that it was power law distribution (Figure 4A and Supplementary Figure 4A). Because the transcriptional regulation network of coding genes is also strictly following power law distribution, this result suggests that TAH-circRNAs share similar transcription mechanism by with coding genes.

Again highly similar as the transcriptional pattern of coding genes (Hnisz et al., 2015; Peeters et al., 2015; Saint-Andre et al., 2016), some of the circRNAs were occupied by a large number of TFs (lower right corner of Figure 4A). It has been well-established that a gene (often termed “Hub gene”) with more interacting partners or sharing more upstream TFs with other genes is potentially more significant functionally in relevant biological processes (Arnone and Davidson, 1997; Wagner, 1999; Odom et al., 2006). Thus, these “Hub circRNAs” may likewise have important functions.

We observed that top-ranked TFs occupying TAH-circRNAs were similar with top-ranked TFs occupying coding genes (Figure 4B and Supplementary Figures 4C–E), suggesting that some master TFs have similar significance for regulation of host...
FIGURE 2 | Epigenomic analyses of TAH-circRNAs in human cell lines. (A) Venn diagram showing the overlap between increased H3K27ac signal intensity in proximal regulatory region of circRNAs. P-value was performed using hypergeometric test. (B) Box plots showing the correlation between H3K27ac signal intensity in proximal regulatory region of circRNAs and RPKM level of circRNAs. P-value was calculated by Pearson correlation. (C) Metagene representation of indicated ChIP-seq occupancy at either TAH-circRNAs or other circRNAs in HepG2 cell line. TSS, transcription start site of circRNAs. (D) Conservation scores of circRNAs in indicated cell lines. (E) IGV plots of RNA-seq and ChIP-seq profiles for the indicated histone marks and TFs for hsa_circ_0061137 and its host gene DIDO1 in HepG2 cell line. (F) The transcriptional activity of TSS (hsa_circ_0061137_upstream 1,500/1,000 nt) was assessed by luciferase reporter assay in HepG2.
genes and circRNAs. Although a number of circRNAs share TFs with their host genes, some TFs were found to only bind to the regulatory region of circRNAs, but not that of their host genes (between 2 Kb upstream and 1 Kb downstream of the host TSS) (Figure 4C). Moreover, relative to other circRNAs, the regulatory regions of TAH-circRNAs had more unique TF binding (Figure 4C). The result imply that a proportion of TAH-circRNAs may be regulated by certain specific TFs independent of host genes.

We found that ~93% TF-circRNA regulatory relationships occurred in only one out of six cell lines (Figure 4D) and only ~1% such regulatory relationships were identified in more than four cell lines. This distribution pattern is again highly consistent with the regulation pattern between TF-coding gene, which has strong cell type- and lineage-specificity (Barshir et al., 2014).

We next asked whether circRNAs occupied by cell-type specific TFs are preferentially expressed in corresponding cells. Since the majority of the ENCODE ChIP-seq data was produced from four cell lines (K562, GM12878, HepG2, and H1-hESC), this analysis was restricted to these four cells. We highlighted several TFs and circRNAs that were highly expressed in a particular cell line. For example, we found that RUNX3 specially bound to the proximal regulatory region of hsa_circ_0088154 in GM12878 cells, and both RUNX3 and hsa_circ_0088154 were only expressed in GM12878 cells (Figure 5). Moreover, proximal regulatory region of hsa_circ_0088154 had more H3K27ac signals in GM12878 than in the other three cell lines (top right inset in Figure 5). In contrast, RGS3, the host gene of hsa_circ_0088154, had a different and inconsistent expression pattern (top right inset in Figure 5). As another example, we found that FOXA1 interacted with the hsa_circ_0004865 only in HepG2 cell line (FOXA1 and hsa_circ_0004865 are shown in the lower left inset of Figure 5). FOXA1 and hsa_circ_0004865, but not its host gene SPTA13, were only highly expressed in HepG2 cells (Supplementary Figure 4B). These results suggest that TAH-circRNAs are under the regulation of specific TFs in a cell type-specific manner.

Characterization of SE-Assigned CircRNAs

Super-enhancers (SEs) play key regulatory roles in transcriptional activation in both normal cell development and disease states (Hnisz et al., 2013; Loven et al., 2013; Jiang et al., 2019; Qian et al., 2019; Bai et al., 2020). Very recently, Bin et al. identified an SE-regulated circRNAs (circNfix), which has important functions in cardiomyocyte proliferation and angiogenesis (Huang et al., 2019). However, it is unknown if this report merely represents a particular case. We thus next systematically analyzed the role of SEs in the transcriptional regulation of circRNAs. Importantly, we found that SE-associated circRNAs were up-regulated...
compared with TE-associated circRNAs in all analyzed cell lines (Figures 6A,C).

Furthermore, the expression levels of SE-associated circRNAs were higher than other circRNAs in HCC clinical samples (one-sided Wilcoxon rank-sum test, P-value < 0.05, Figure 6B). Consistently, the regulatory regions of SE-associated circRNAs have more TF binding (Figure 6D) and higher H3K4me1 signals (Supplementary Figure 5C) than TE-associated circRNAs. We confirmed the expression of randomly selected twelve circRNAs associated with SEs by Real-time PCR and Sanger sequencing (Table 1, Supplementary Figures 5A,B).

One exemplary SE-associated circRNA (has_circ_0001827) was shown in Figure 6E. A prominent SE cluster (Red bar in Figure 6E) was identified upstream of this circRNA, which also had notable H3K36me3 signals along its gene body, indicative of active transcriptional elongation. Importantly, ChIA-PET results identified that the TSS of has_circ_0001727 had direct DNA-DNA contact with three SE constituents (Shaded regions in Figure 6E). These SE constituents were co-occupied by several TFs, including POLR2A, HNF4G, CEBPB, RXRA, and USF1. Moreover, the proximal regulatory region of this circRNA was directly occupied by RXRA and USF1. Lastly, compared with its host gene, has_circ_0001727 had much higher expression levels (>17-fold). These results together strongly suggest that has_circ_0001727 is transcriptionally activated by several TFs via its upstream SE region. In addition, we found that 38–70% TAH-circRNAs were associated with TEs, and 2–10% among these circRNAs were associated with SE (Figure 6F).

**Identification and Validation of TAH-CircRNAs Transcriptionally Regulated by FOXA1**

Our earlier results (Figure 4B) showed that FOXA1 had the highest number of binding sites with in regulatory regions of TAH-circRNAs. We thus next sought to functionally validate the transcriptional regulation on TAH-circRNAs, using FOXA1 as a model TF.

To this end, we silenced FOXA1 using siRNA in HepG2 cells and quantified the genome-wide changes of the expression
of both coding genes and circRNAs using microarrays. As a result, we identified a total of 2,136 circRNAs downregulated (FC < 1.4) following FOXA1 silencing. Importantly, 65 of FOXA1-occupied TAH-circRNAs (total n = 1,201) were significantly over-presented in these down-regulated circRNAs (Figure 7A). We further performed permutation test by sampling 20,000 times random circRNA sets and confirmed that the over-presentation of FOXA1-occupied TAH-circRNAs was not by chance (Figure 7B). Moreover, for the overlapped 65 circRNAs, the expressions of their host genes were not altered by FOXA1-silencing (Figure 7E). In addition, Gene set enrichment analysis (GSEA) demonstrated that FOXA1-occupied TAH-circRNAs were significantly enriched in the down-regulated transcripts upon knocking down of FOXA1 (Figure 7D). In contrast, this significance was not observed in the host genes of these circRNAs (Figure 7C).

Figure 7F shows an example of FOXA1-occupied TAH-circRNA (hsa_circ_0020306) whose expression was decreased following FOXA1 knockdown (Fold change = 0.831). Notably, ChIA-PET identified direct interactions between the TSS of this circRNA and distal regions with both H3K27ac and H3K4me1 signals (indicative of potential enhancers). Moreover, FOXA1 directly occupied both the TSS and distal regions together with other TFs including Pol II, strongly suggesting a direct transcriptional regulation of this circRNA. In contrast, expression of its host gene (CHST15) was upregulated after knocking down of FOXA1 (1.062-fold). Taken together, these results suggested that a set of TAH-circRNAs are under direct regulation by FOXA1 independent of their host genes.
CircRNAs are increasingly recognized as important players in many biological processes (Legnini et al., 2017; Tang et al., 2019). For example, circRNAs are capable of regulating gene expression by competing for protein and miRNA binding (Hansen et al., 2013; Du et al., 2016). They also compete with linear RNA production and regulate the accumulation of full-length mRNA (Ashwal-Fluss et al., 2014). However, upstream regulatory mechanisms of the transcription of circRNAs are still largely unknown. Here, we performed comprehensive epigenomic investigations of human circRNAs, analyzing transcriptome, cistrome, and DNA methylome data from a total of 91 samples. We identified a group of TAH-circRNAs with distinct epigenomic and biological features, including (i) the regulatory region of TAH-circRNAs...
TABLE 1 | These circRNAs were verified by Real-time PCR and they are associated with SE.

| CircRNA_ID | CircRNA_name | Chr | Start    | End     | Fold_change | Hostgene |
|------------|--------------|-----|----------|---------|-------------|----------|
| circ_4565  | hsa_circ_0004565 | 3   | 149663797| 149619949| 3.358573    | RNF13    |
| circ_8301  | hsa_circ_0008301 | 11  | 1307231  | 1317024 | 5.081821    | TOLLIP   |
| circ_0515  | hsa_circ_0000515 | 14  | 20811305 | 20811534 | 2.885618    | RFFH1    |
| circ_4004  | hsa_circ_0004004 | 5   | 17239438 | 17236213 | 2.853213    | ERGIC1   |
| circ_7099  | hsa_circ_0007099 | 15  | 89656955 | 89659752 | 2.12391     | ABHD2    |
| circ_6913  | hsa_circ_0006913 | 1   | 27087346 | 27089776 | 3.229994    | ARID1A   |
| circ_5188  | hsa_circ_0005188 | 19  | 47858295 | 47858607 | 6.494322    | DHX34    |
| circ_9392  | hsa_circ_00049392 | 19 | 11226769 | 11227674 | 2.137966    | LDLR     |
| circ_3006  | hsa_circ_0003006 | 3   | 58111307 | 58112489 | 6.272582    | FLNB     |
| circ_7769  | hsa_circ_0007769 | 6   | 14408693 | 144086935| 3.981750    | PHACTR2  |
| circ_0777  | hsa_circ_00040777 | 16 | 87453388 | 87454322 | 2.029921    | ZCCHC14  |
| circ_7361  | hsa_circ_0007361 | 16  | 87451065 | 87452507 | 2.094044    | ZCCHC14  |
| circ_1727  | hsa_circ_0001727 | 7   | 99621041 | 99621930 | 1.681775    | ZKSCAN1  |

FIGURE 7 | Identification and validation of TAH-circRNAs transcriptionally regulated by FOXA1. (A) Venn diagram showing the overlap between the down-regulated circRNAs after FOXA1 knockdown (Left) and circRNAs bound by FOXA1 (Right). P-value was performed using hypergeometric test, ***P < 0.001. (B) Gene set enrichment analysis of the expression changes of FOXA1-binding TAH-circRNAs and (C) their corresponding host genes. (D) The number of down-regulated circRNAs overlapping with the TAH-circRNAs that were bound by FOXA1 (black bar) compared with 20,000 random circRNA sets (gray bars). R1: randomly sampling from 92,375 human circRNAs from circBase database. R2: randomly sampling from 19,805 human circRNAs that were bound by FOXA1. (E) Expression changes of the 65 FOXA1-binding TAH-circRNA (left) and their corresponding host genes (right) in response to FOXA1 knockdown. NC, negative control. (F) IGV plots of RNA-seq and ChIP-seq profiles for the indicated histone marks and TFs for hsa_circ_00203006 and its host gene CHST15 loci in HepG2 cells.
had higher signals of H3K36me3 and DNaseI than other circRNAs and, consistently, TAH-circRNAs were occupied by more TFs, suggesting stronger transcriptional activation; (ii) TAH-circRNAs had greater conservation score than other circRNAs, indicating their function significance; (iii) the regulatory regions of TAH-circRNAs are markedly more hypo-methylated than other circRNAs, in agreement with its higher transcriptional output.

Most importantly, we provided a number of lines of evidence to demonstrate that TAH-circRNAs are under direct transcriptional regulations mediated by TFs. First, ChIA-PET results identified that 23–52% of TSS of TAH-circRNAs had direct interactions with distal cis-regulatory regions. In comparison, only 11–46% of non-TAH-circRNAs displayed such contacts. As shown in our exemplary circRNAs (Figures 2E, 6E, 7F), these DNA-DNA interactions almost always occurred between the circRNA TSS and potential enhancer regions (positive of enhancer signals and TF binding), strongly suggesting that these TAH-circRNAs can be directly transcribed independent of host genes. Second, we found a subset of TF-circRNA regulatory pairs that did not involve their host genes (Figure 5). Third, we identified SE-associated circRNAs that had higher expression levels than other circRNAs. Fourth, functional validation further confirmed that FOXA1 directly regulated the transcription of at least 65 TAH-circRNAs independent of their host genes.

In summary, this work systematically analyzes the epigenomic features of circRNAs and identifies a distinct group of TAH-circRNAs. Moreover, our study demonstrates that independent transcriptional mechanisms exist for the regulation of the expression of circRNAs. These results together shed important insights into the regulation of this large class of non-coding RNAs.

REFERENCES

Arnone, M. I., and Davidson, E. H. (1997). The hardwiring of development: organization and function of genomic regulatory systems. Development 124, 1851–1864. 
Ashwal-Fluss, R., Meyer, M., Pamudurti, N. R., Ivanov, A., Bartok, O., Hanan, M., et al. (2014). circRNA biogenesis competes with pre-mRNA splicing. Mol. Cell 56, 55–66. doi: 10.1016/j.molcel.2014.08.019 
Bai, X., Shi, S., Ai, B., Jiang, Y., Liu, Y., Han, X., et al. (2020). ENDb: a manually curated database of experimentally supported enhancers for human and mouse. Nucleic Acids Res. 48, D51–D57. doi: 10.1093/nar/gkz973 
Barshter, R., Shwartz, O., Smoly, I. Y., and Yeger-Lotem, E. (2014). Comparative analysis of human tissue interactomes reveals factors leading to tissuespecific manifestation of hereditary diseases. PLoS Comput. Biol. 10:e1003632. doi: 10.1371/journal.pcbi.1003632 
Chen, L. L. (2016). The biogenesis and emerging roles of circular RNAs. Nat. Rev. Mol. Cell Biol. 17, 205–211. doi: 10.1038/nrm.2015.32 
Dong, R., Ma, X. K., Chen, L. L., and Yang, L. (2017). Increased complexity of circRNA expression during species evolution. RNA Biol. 14, 1064–1074. doi: 10.1080/15476286.2016.1269999 
Du, W. W., Wang, Y., Liu, E., Yang, Z., Dhaliwal, P., and Yang, B. R. (2016). Foxo3 circular RNA retards cell cycle progression via forming ternary complexes with p21 and CDK2. Nucleic Acids Res. 44, 2846–2850. doi: 10.1093/nar/gkw027 

DATA AVAILABILITY STATEMENT

The accession number for the microarray data reported in this paper is NCBI GEO: GSE161174.

AUTHOR CONTRIBUTIONS

X-CL and Z-DT analyzed data and wrote the paper. Y-YL, F-CQ, J-MZ, ML, JZha, X-FB, JZhu, and C-CF participated in the pre-processing of the datasets and performed the computational analysis. LP, I-WD, and X-JD performed the experiment validation. Q-YW, JP, and C-QL conceived the idea for the paper, provided the guidance, and critically revised the paper. All authors read and approved the final version to be published.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2020.590672/full#supplementary-material

Feng, C., Song, C., Liu, Y., Qian, F., Gao, Y., Ning, Z., et al. (2020). KnockTF: a comprehensive human gene expression profile database with knockdown/knockout of transcription factors. Nucleic Acids Res. 48, D93–D100. doi: 10.1093/nar/gkz881 
Hansen, T. B., Jensen, T. I., Clausen, B. H., Bramsen, B. J., Finsen, B., Damgaard, C. K., et al. (2013). Natural RNA circles function as efficient microRNA sponges. Nature 495, 384–388. doi: 10.1038/nature12199 
Hnisz, D., Abraham, B. J., Lee, T. I., Lau, A., Saint-Andre, V., Sigova, A. A., et al. (2013). Super-enhancers in the control of cell identity and disease. Cell 155, 934–947. doi: 10.1016/j.cell.2013.09.053 
Hnisz, D., Schuiers, J., Lin, C. Y., Weintraub, A. S., Abraham, B. J., Lee, T. I., et al. (2015). Convergence of developmental and oncogenic signaling pathways at transcriptional super-enhancers. Mol. Cell 58, 362–370. doi: 10.1016/j.molcel.2015.02.014 
Hsiao, K. Y., Lin, C. C., Gupta, S. K., Chang, N., Yen, L., Sun, H. S., et al. (2017). Noncoding effects of circular RNA CCDC66 promote colon cancer growth and metastasis. Cancer Res. 77, 2339–2350. doi: 10.1158/0008-5472.CAN-16-1883 
Huang, S., Li, X., Zheng, H., Si, X., Li, B., Wei, G., et al. (2019). Loss of super-enhancer-regulated circRNA Nfix induces cardiac regeneration after myocardial infarction in adult mice. Circulation 139, 2857–2876. doi: 10.1161/CIRCULATIONAHA.118.038361 
Jiang, Y., Qian, F., Bai, X., Liu, Y., Wang, Q., Ai, B., et al. (2019). SEdb: a comprehensive human super-enhancer database. Nucleic Acids Res. 47, D235–D243. doi: 10.1093/nar/gky1025
