Heterogeneous circRNA expression profiles and regulatory functions among HEK293T single cells

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The single-cell analysis is becoming a powerful method for early detection of the abnormal variant in tissues, especially for profiling a small number of heterogeneous cells. With the advancement of sequencing technologies, many types of non-coding elements including miRNAs and lncRNAs which shed light on their heterogeneous patterns and functions among cells, have been profiled at the single-cell level. However, the complete picture of circRNA profile at single-cell level is still lacking. In this study, RNA-Seq data obtained from single HEK293T cells have been used to analyze expressions and functions of heterogeneous circRNA profiles. The enrichment patterns of circRNAs, interactions with miRNAs and pathways such as ErbB signaling pathway and protein processing in endoplasmic reticulum, have also been investigated. The results showed that circRNAs had a specific distribution pattern which was implicated with expression, miRNA and functional profiles at single-cell level. This assessment study of the expressions and functions of circRNAs at single-cell level shed light on heterogeneities among single cells.

In recent years, with the advancement of next generation sequencing technologies, great progress has been made in transcriptome researches, most of which were paid attention to bulk samples. However, intrinsic heterogeneity has been identified to be widespread within the transcriptomes of different individual cells, even within the same types of cells. Moreover, previous investigations showed that these genetic heterogeneities might be averaged out in bulk sequencing, especially for those rare non-coding RNAs who are dynamically expressed in cells. The single-cell technology can profile heterogeneities within the same tissue at the single-cell level and serve as a powerful method to identify specific properties of each cell. Due to the superiority in detecting single-cell heterogeneity, single-cell technology has become the focus in many fields. And a variety of single-cell sequencing methods, such as CEL-seq, Quartz-Seq, Smart-seq, MATQ-seq, make the detection of transcriptional variation in single cells accessible and meet the demand for anatomical resolution. In particular, the study of non-coding RNA at single-cell level has attracted extensive attention.

Circular RNA (circRNA) is one of the new members of the non-coding RNA family, forms in a covalently closed continuous loop and isn’t terminated at 5' and 3' ends. With the development of high-throughput RNA sequencing technology, abundant circRNAs have recently been identified in many kinds of species and implicated in important functions in physiological and disease process. Hence, several comprehensive databases such as CircBase, Circ2Traits, circRNABase, deepBase, and circRNAdb, have been developed to merge and unify information of published circRNAs and provide a series of online alignment tools to maintain structure and function prediction. To meet the demand of comprehensive detection of circRNAs, several different pipelines with better performance in circRNA analysis have been developed, including find_circ, CIRI, circRNAFinder, CIRCexplorer, UROBORUS, etc., for further study of circRNAs. However, most of existing work on circRNA is based on bulk sequencing, in which individual cell properties and heterogeneities are hidden. Furthermore, although so much progress has been made in circRNA analysis, neither the circRNA expression pattern nor the specific circRNA function at the single-cell level has been reported. CircRNA analysis at single cell level can yield more detailed and accurate genetic information, and provide a clue for dynamic variation of circRNAs and a new

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In this study, to get a high-resolution profile of circRNAs for describing the distribution at the single-cell level, single-cell sequencing datasets were collected. Firstly, we performed expression profile and heterogeneity pattern analysis at single-cell level. Secondly, we elucidated the correlation between the expression of circRNAs and their host genes. Thirdly, by implementing GO analysis, we were interested in the functional enrichment and pathways of the circRNA host genes. Furthermore, we examined the potential function of miRNAs as sponges by inference of circRNA regulatory networks. Our data provided a novel basis for circRNA research in single cells. It was found that the single-cell circRNA profiles might have a specific distribution pattern which was implicated with expression, miRNA and functional profiles at the single-cell level.

Results

Detection results of general circRNA by using different methods. We obtained four sets of detection results generated by CIRI2, circRNAFinder, find_circ and CIRCexplorer, respectively. The number of circRNAs ranged from 1,111 to 6,493 in 38 single cells, in which only 410 circRNAs were predicted by all the four methods. In addition, we also detected 68 circRNAs in 7 another single cells from GSE53386. To avoid the technical bias, we focused on the 410 circRNAs found by all the four methods, and also the 68 circRNAs in 7 another single cells were taken into account.

Among circRNAs, heterogeneity was found in each cell. The number of circRNAs in these single cells was significantly different (t-test, \( p = 7.338e-15 \)), and the types of circRNAs were also quite different. To characterize the level of heterogeneity, Manhattan distance was used to calculate the distances from the pairwise single cells (Supplementary Fig. 1A). Moreover, heterogeneity was indicated by the hierarchical comparison in each cell and hierarchical clustering showed that heterogeneities of circRNA expression were distinct among cells. In particular, single-cell samples such as SRR5091997 and SRR5091976, which had the similar quantity of sequencing reads, were quite different in circRNAs. Due to the differences of samples, the correlation of samples was analyzed and the hierarchical clustering method was applied to group circRNAs. Samples had only weak correlation or the correlation of samples was analyzed and the possible correlation between SNP-Freq and circRNA-Freq, and the result suggested that there was no linear correlation between them (Fig. 2A). The similar SNP profiles and correlation were also found in the 7 single cells from GSE53386 (Supplementary Fig. 2A).

Comparisons of circRNA and SNP profiles of single cells. To find an overview of the SNP distribution and the possible enrichment of SNPs on chromosomes, the SNP frequency on each chromosome was normalized by the quantity of the SNPs and the length of chromosomes (SNP-Freq, refer to Formula (2) in the Materials and Methods), the distribution of the circRNAs on the chromosomes was depicted based on the results from the 38 single cells (Fig. 1A). The Heatmap showed that the distribution of circRNAs on chromosomes was not uniform and enriched on chr22 (\( p < 2.86e-24 \)). In addition, cellular heterogeneity was found among samples, although the cells considered here were cultured in the same condition. The same result was found in 7 single cells from GSE53386 (Supplementary Fig. 2A).

Functions of circRNA host genes in single cells. Gene Ontology annotation can evaluate the function enrichment, as well as to gain an insight into functions of all genes harboring circRNAs. Each gene was associated
with at least one GO term and had a wide range of biological functions. According to three categories: biological processes (BP), molecular function (MF) and cellular component (CC), all of GO terms were classified to point out the significantly overlapped functions. In the BP, the circRNA host genes were significantly enriched transcription, DNA–templated, regulation of transcription DNA–templated (p < 1.04e-20) (Fig. 3A). In the MF, genes were significantly enriched in protein binding (p < 5.56e-31) (Fig. 3B). In the CC, genes were significantly enriched in nucleus, nucleoplasm and cytoplasm (p < 1.97e-28) (Fig. 3C). These GO terms associated with circRNA host genes were significantly correlated with the transcription, protein binding, etc., which indicated that

Figure 1. Heatmap of circRNA distribution on the chromosomes in 38 single cells. (A) The circRNA enrichment on chromosomes for the 38 single-cell samples; (B) Heatmap of the SNP-Freq on each chromosome for 38 single-cell samples obtained by the GATK and Samtools.
those genes were more likely to undergo circularization, and circRNAs might be crucial in the regulation of these functions during embryonic development.

**Pathway analysis of circRNA host genes.** Genes harboring circRNAs were observed to be involved in different pathways (Table 1), even some of which had a high "sample ratio" (a ratio of samples with at least one circRNA to the total sample). The circRNA host genes $PIK3CB$, $PTK2$, $CRKL$, $PIK3R3$, $BRAF$, $ABL2$ and
MTOR (Fig. 4A) were assigned to the ErbB signaling pathway, and many of them were linked to the enriched GO terms such as protein binding and nucleolus. The ErbB signaling pathway, which regulated diverse physiological responses such as cell survival, proliferation and motility. This consistency between the GO terms and the genes might indicate the possible regulation of these functions in cell metabolism. CRKL, which regulated cell adhesion, spreading and migration, harbored circRNAs in 12 cells. And CRKL harboring SNP was found in 22 cells. Upon testing the relationship between CRKL expression among the cells, we found that the expression of CRKL...
increased (t-test, \( p = 0.008 \)) in cells with circRNAs. And also checked the expression of CRKL in cells with SNPs and no significant differences existed among the cells. Whether SNPs affected the expression of CRKL or not was needed to be further studied.

Another pathway protein processing in endoplasmic reticulum, contained circRNA target genes PDA6, SEC, 31B, UBE4B, UBQLN1, UBE2G1, SEC, 62, NPOC4, AMFR, and MAN1A2 (Fig. 4B). These genes which played an important role in protein folding, translocation, and degradation, had varying degrees of sample ratios. Among these genes, MAN1A2 which had Alpha-mannosidases function during the N-glycan maturation process also harbored circRNAs in 12 cells with comparatively higher expression. This circularization in checkpoint genes might have potential effects on their expression regulation and was likely to be important factors in maintaining normal physiological function.

**CircRNA-miRNA-mRNA associations.** Interactions, which indicate the functions of circRNAs, between circRNAs and their target miRNAs were predicted according to complementary seed sequence matches. A total of 249 miRNAs could be combined with 406 circRNAs. Among them, the hsa-miR-15a, hsa-miR-15b, hsa-miR-16, hsa-miR-195, hsa-miR-424 and hsa-miR-497 regulated the largest number of circRNAs. Furthermore, investigations on the associations between each potential complementary binding miRNA and their target genes of human diseases were performed. Among the potential complementary binding miRNAs of circRNAs, 120 miRNAs could be associated with a broad spectrum of diseases by acting on 305 disease-related genes. Thus, an entire network of circRNA-miRNA-mRNA interactions was delineated by using Cytoscape (Fig. 5A).

Further, the potential relevance of circRNA and mRNA was established by miRNA to analyze the molecular mechanism. It was predicted that circMAN1A2 could harbor hsa-miR-494, hsa-miR-491-5p, hsa-miR-433, hsa-miR-384, hsa-miR-543, hsa-miR-107, hsa-miR-301b, hsa-miR-329, hsa-miR-152, and hsa-miR-362-3p by miRNAs seed sequence matching, respectively. Moreover, DIANA-miRPath analysis revealed that a total of genes could be regulated by these ten potential miRNAs (Supplementary Table S1), and these target genes were associated with ErbB signaling pathway. As a result, a network of circRNA-miRNA-mRNA interactions in ErbB signaling pathway was established (Fig. 5B).

ErbB signaling pathway network containing 33 genes on circMAN1A2 mediated by hsa-miR-494, hsa-miR-491-5p, hsa-miR-433, hsa-miR-384, hsa-miR-543, hsa-miR-107, hsa-miR-301b, hsa-miR-329, hsa-miR-152, and hsa-miR-362-3p was also established (Fig. 6A). TGFA could be targeted by hsa-miR-384, hsa-miR-543, hsa-miR-301b, hsa-miR-329, hsa-miR-152 and hsa-miR-362-3p in the network, which mean that it might be a crucial factor mediated by circMAN1A2. These findings suggested that circMAN1A2 should participate in the ErbB signaling pathway. When we linked these targeted genes to their GO terms, the GO annotation for target genes revealed that genes might indicate the possible regulation of these functions in embryonic development at single cells.

**Discussions**

In the past few years, circRNA was identified to be highly abundant in mammals based on the study of population cells. However, analysis of population cells could only detect the characteristics of the average, which limited to obtain clues to the differences among each cell. Fortunately, the advent and utilization of single-cell sequencing facilitated the research on heterogeneity and functions of circRNA. In an analysis of publicly available RNA-seq data from HEK293T single cells, we found that circRNAs were dynamically expressed in single cells.

In this study, we reported the heterogeneity patterns of circRNAs at the single-cell level as well as gene expressions. We obtained 410 circRNAs in 38 single cells, and found that cell-level heterogeneity of circRNAs was obvious, which indicated that circRNA exhibited different expression patterns. These data provided a more comprehensive view into circRNAs than that in published reports in bulk sequencing. The distribution on chromosomes profiles revealed circRNAs were distributed on all chromosomes except for Y chromosome. This distribution was not uniform on the chromosomes and got an enrichment on chr22. And SNPs were widely distributed on 24 chromosomes and were enriched on chr16, chr17, chr19 and chr22. Especially, the possible correlation of circRNA-Freq and SNP-Freq suggested that no correlation between them though SNPs was detected in 166 circRNA host genes. Whether they were relevant or not was still to be further studied. The expression of circRNA had recently been

| Pathway | circRNA host genes |
|---------|--------------------|
| Lysine degradation** | HADH, ASH1L, SETD2, WHSC1, NSD1, WHSC1L1, EHMT1 |
| Ubiquitin mediated proteolysis** | HUWE1, SMURF1, TRIM57, UBE4B, UBE2G1, CUL5, BIRC6, UBE2I, UBR5, MID1 |
| ErbB signaling pathway** | PIK3CB, PT2, CRKL, PIK3R3, BRAF, ABL2, MTOR |
| Protein processing in endoplasmic reticulum** | PDI, SEC, UBE4B, UBQLN1, UBE2G1, SEC, 62, NPOC4, AMFR, MAN1A2 |
| Insulin signaling pathway** | PIK3CB, CRKL, PIK3R3, BRAF, ACACA, ACACA, PRKAK2, MTOR |
| RNA transport* | NUP214, RANGAP1, KPNB1, CASC3, UBE2I, EIF4G3, NUP205, NDC1 |
| Amoebiasis* | PIK3CB, PT2, PLCB1, PIK3R3, ADCY1, RAB7A |
| Selenocompound metabolism* | MTR, PAPS1, MARS |
| Bacterial invasion of epithelial cells* | PIK3CB, PT2, CRKL, PIK3R3, CD2AP |
| AMPK signaling pathway* | PIK3CB, PIK3R3, ACACA, PRKAK2, PPARG, MTOR |

Table 1. Representative circRNA host genes in the pathway analysis. Note: Genes harboring circRNAs were observed to be involved in ten pathways by p-value cutoff at 0.05. (*p < 0.05, **p < 0.01).
The Pearson correlation coefficient was calculated between the expression levels of circRNAs and their host genes. The results showed a strongly negative correlation between circRNAs and their host mRNA, suggesting that these circRNAs have potential influence on the transcription of their host genes. Meanwhile, SNP heterogeneity was also found in 7 single cells from GSE53386.

GSEA was used to reflect enrichment situation of circRNA host genes in three ranked gene lists, including all expressed genes, all genes harboring SNPs and miRNAs. GSEA walked down the ranked list, and the enrichment score was increased if the gene was present in a gene set, otherwise, decreased. The magnitude of the increase or decrease was determined by the correlation of genes with expression level, SNPs and miRNAs. For both GSE78968 and GSE53386 datasets, there were three patterns of circRNAs: (i) the expression of circRNA host genes was higher than those without circRNA; (ii) random distribution in SNPs of the circRNA host genes; (iii) no expression of circRNA host genes.

Figure 4. ErbB signaling pathway and its circRNA-related genes. (A) The ErbB signaling pathway and its gene interactions; (B) Protein processing in endoplasmic reticulum and its gene interactions. The light green boxes represent circRNA-free genes among all single cells, the yellow boxes represent genes harboring circRNAs appearing in only a single cell and the orange boxes represent genes harboring circRNAs appearing in multi-samples.

identifying a correlation between circRNAs and linear host mRNA, and even for regulate transcription of their host genes. In the present study, we had calculated the Pearson correlation coefficient between the expression levels of circRNAs and their host genes. The results showed that a strongly negative correlation between circRNAs and their host mRNA existed, which suggested that those circRNAs have potential influence on the transcription of their host genes. Meanwhile, SNP heterogeneity was also found in 7 single cells from GSE53386.

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observed; (iii) circRNA host genes contained more targeted miRNAs. The common features of the tendency in expression level and miRNA counts had a potential impact on RNA circularization. Therefore, we might conclude that circRNAs might function as miRNA sponges to improve target host gene expression.

Results from GO function annotation and KEGG pathway help to enrich and identify important genes generating circRNAs. GO function annotations for those target genes revealed that they were significantly implicated in protein binding, nucleus, nucleoplasm and cytoplasm, which provided evidence for embryonic development-related circRNA accumulation. Considering KEGG pathway, those circRNA host genes involved important physiological pathways such as ErbB signaling pathway, and protein processing in endoplasmic reticulum, which reflected their importance in embryonic development. Circularization in these checkpoint genes might have potential effects on the ErbB signaling pathway and protein processing in endoplasmic reticulum, which suggested that circRNAs be likely to play regulatory roles in development. As stated in the published study, circRNAs could be correlated with regulation mechanisms of embryonic development.

In addition, we discovered that circRNAs harbored substantially miRNA target sites based on conservative seed sequence matches. Hundreds of circRNA-miRNA interactions were predicted, which would supply new discernment for the underlying mechanisms. CircRNAs could be correlated with disease miRNAs and the circRNA-miRNA axes might participate in disease-related pathways. The circRNA-miRNA-mRNA network could serve as the powerful regulation pathway for the cascade amplification effect of circRNA-miRNA and miRNA-mRNA. The best-known biological impact of circRNAs was playing miRNA sponge effects. Therefore, we speculated that circMAN1A2 might competitively bind with hsa-miR-494, hsa-miR-491-5p, hsa-miR-433, hsa-miR-384, hsa-miR-543, hsa-miR-107, hsa-miR-301b, hsa-miR-329, hsa-miR-152, and hsa-miR-362-3p, and had effects on associated target genes. DIANA-miRPath determined the candidate miRNAs of circMAN1A2 which were involved in the ErbB signaling pathway. These miRNA target genes were enriched for functional annotations relating to biological regulation, protein binding, as well as development. Previous studies had revealed that abnormal ErbB signaling in humans was associated with the development of neurodegenerative diseases or a wide variety of types of solid tumor. Thus, we predicted that circMAN1A2 could act as a regulator of the ErbB signaling pathway.

Figure 5. circRNA-miRNA-mRNA networks. (A) The circRNA-miRNA-mRNA network, which consists of 406 circRNAs (orange nodes), 249 miRNAs (blue nodes) and 305 disease genes (green); (B) The prediction of circMAN1A2 target genes related to ErbB signaling pathway. A total of 44 nodes and 72 edges were constructed.

CircRNA within single-cells represented one type of the non-coding elements among heterogeneous cells, and we believed that more interesting finding for non-coding element analysis might be revealed based on more complete and accurate single-cell sequencing data. Thus, it was quite promising that the single-cell sequencing was allowing scientists to explore non-coding small molecules diversity in cell populations.

Conclusion
In this single-cell RNA-Seq analysis of HEK293T cells, not only the circRNA distribution patterns but also the single-cell SNP, gene expression and function profile were profiled. It was found that circRNAs were dynamically expressed in single cells and had obvious heterogeneity by the analysis of distribution patterns. These circRNAs were potentially involved in transcription, DNA-templated, protein binding, and nucleus. Moreover, these circRNAs had a specific distribution pattern which was not associated with SNPs but was implicated with gene expressions or functional profiles at the single-cell level. In addition, the circMAN1A2 might serve as a regulator of the ErbB signaling pathway. These data laid a foundation for further decipher characteristics and regulation mechanisms of circRNAs in single cells.
Materials and Methods

RNA-Seq data for the single-cell samples. Two data sets of single HEK293T cells were downloaded from NCBI GEO database (Supplementary Table S2), the first data set (GEO ID: GSE78968) had 38 single cells, which were all obtained from the same HEK293T clone and performed by MATQ-seq. Second data set (GEO ID: GSE53386) had 7 single HEK293T cells, which were performed by SUPeR-seq. Reads of both two data sets were used for the following analyses of circRNA and SNP.

CircRNA detection. Quality filtering on these reads was performed using Parallel-QC and sequences which did not fulfill the following criteria were discarded: reads with quality score ≥ 20, the GC proportion ranged from 0.4 to 0.6. All the downstream analyses were based on clean data.
The reference genome and annotation used in the following analyses were downloaded from UCSC Genome Browser (http://genome.ucsc.edu) (hg19 version). Four pipelines, including CIRI, version 2.0.1, circRNAFinder, version 0.1.0, find_circ and CIRCexplorer, version 1.1.10, were introduced to detect the circRNA for the sake of higher accuracy and sensitivity since no specific circRNA detection method has been customized for single-cell RNA-Seq data. All the four pipelines were used the default parameters.

Manhattan distance calculation. To calculate the degree of heterogeneity in each cell, the Manhattan distance was based on the absence or presence of each circRNA. For each cell, a full (all circRNAs by all cells) 0/1 matrix was built, with “1” denoting presence (defined as the detection of circRNA) and “0” denoting no presence of the corresponding circRNA. Thus, Manhattan distance was used to calculate the distance from the matrix.

CircRNA enrichment on chromosomes. For circRNA enrichment on chromosomes, the circRNA frequency on each chromosome was normalized by the following formula:

\[
\text{circRNA} - \text{Freq} = 10^6 \times \frac{\text{circRNA counts/Chromosome}}{\text{chromosome length}}
\]  

Where circRNA counts/Chromosome was the quantity of the circRNAs detected on one chromosome, and chromosome length was the length of this chromosome. The factor $10^6$ was chosen as the denominator to leverage the circRNA-Freq values for a fair and easy comparison.

SNP identification. Reads were mapped to the human genome using BWA software, version 0.7.15. The variants calling was performed in GATK and Samtools, version 1.3. HaftlypeCaller was used for variant calling in GATK while mpileup and view were used in Samtools. The overlaps of GATK and Samtools results were considered as candidates, which were subjected to additional filtering to remove SNPs with low-quality value (QUAL < 30), low QD (QD < 20.0), low read coverage (DP < 8) and Strand Bias (FS > 30.0).

For SNP enrichment on chromosomes, the SNP frequency on each chromosome was normalized using the following formula:

\[
\text{SNP} - \text{Freq} = 10^6 \times \frac{\text{SNP counts/Chromosome}}{\text{chromosome length}}
\]  

Where SNP counts/Chromosome was the quantity of the SNPs detected on one chromosome, and chromosome length was the length of this chromosome. The factor $10^6$ was chosen as the denominator to leverage the SNP-Freq values for a fair and easy comparison.

Expression analysis. RNA-seq libraries were mapped to the reference genome using tophat, version 2.1.0, after that the FPKM values for each gene were calculated by cufflinks, version 2.2.0 and reads mapped to genomic features were counted using htseq-count. Therefore, to estimate abundance of circRNAs, the circRNA-gene ratio (CGR) was quantified using the following formula:

\[
\text{CGR(circRNA)} = \frac{junction \ spanning \ reads}{host \ gene \ reads}
\]  

To generate an overview of circRNA expression profiles among the single cells, the hierarchical clustering analysis was performed based on expression value of all target circRNAs. Expression levels of circRNAs were quantified by the number of junction spanning reads. To obtain an estimate of relative expression, the number was normalized to the total number of reads in the library and the host gene FPKM. The FPKM of circRNA was calculated using the following formula:

\[
\text{FPKM (circRNA)} = \frac{junction \ spanning \ reads}{mapped \ reads \ (millions) \times \ host \ gene \ FPKM}
\]  

Where junction spanning reads was the amount of the circRNAs detected on one site, and million mapped reads were the total reads which mapped to reference genomes.

miRNA analysis. For miRNA analysis, microRNA target prediction in human was available on Miranda database (http://www.microrna.org/microrna/home.do), which was downloaded to evaluate all instances of conservative sites in each gene. The miRNA-disease associations were predicted in the Human MiRNA Disease Database (HMDD http://cmbi.bjmu.edu.cn/hmdd). All mature miRNA sequences were downloaded from miRBase (http://www.mirbase.org), each circRNA was scanned to identify miRNA target sites based on conserved seed sequence matches. The miRNA pathway investigating was carried out based on DIANA-miRPath by \( p \)-value cutoff at 0.05. The graphs of the circRNA-miRNA and circRNA-miRNA-gene networks were visualized on Cytoscape, version 3.3.0.

Functional enrichment analysis. The function of those target genes was predicted and annotated in the network by Database for Annotation, Visualization and Integrated Discovery (DAVID https://david.ncifcrf.gov), \( P < 0.05 \) was used as the criterion for statistical significance. Gene Ontology (GO) that describes genes from any organism were used. GO Terms were classified into three categories: biological processes (BP), cellular
References
1. Khan, S. et al. Comparative transcriptome analysis reveals candidate genes for the biosynthesis of natural insecticide in Tanacetum cinerariifolium. BMC Genomics 18(1), p. 54 (2017).
2. van Dijk, E. L. et al. Ten years of next-generation sequencing technology. Trends Genet 30(9), p. 418–26 (2014).
3. Cortes-Lopez, M. & Miura, P. Emerging Functions of Circular RNAs. Yale J Biol Med 89(4), p. 527–537 (2016).
4. Shalek, A. K. et al. Single-cell transcriptomics reveals bimodality in expression and splicing in immune cells. Nature 498(7453), p. 236–40 (2013).
5. Graf, T. & Stadtfeld, M. Heterogeneity of embryonic and adult stem cells. Cell Stem Cell 3(5), p. 480–3 (2008).
6. Navin, N. E. Cancer genomics: one cell at a time. Genome Biol 15(8), p. 452 (2014).
7. Lu, W. & Fuchou, T. Recent progress in single-cell RNA-Seq analysis. Yi Chuan 36(11), p. 1069–76 (2014).
8. Fan, X. et al. Single-cell RNA-seq transcriptional analysis of linear and circular RNAs in mouse preimplantation embryos. Genome Biol 16(1), p. 148 (2015).
9. Wang, D. & Bodovitz, S. Single cell analysis: the new frontier in ‘omics. Trends Biotechnol 28(6), p. 281–90 (2010).
10. Navin, N. et al. Tumour evolution inferred by single-cell sequencing. Nature 472(7341), p. 90–4 (2011).
11. Eberwine, J. et al. The promise of single-cell sequencing. Nat Meth 11(1), p. 25–7 (2014).
12. Hashimshony, T. et al. CEL-Seq: single-cell RNA-Seq by multiplexed linear amplification. Cell Rep 2(3), p. 666–73 (2012).
13. Sasagawa, Y. et al. Quartz-Seq: a highly reproducible and sensitive single-cell RNA sequencing method, reveals non-genetic expression heterogeneity. Genome Biol 14(4), p. R31 (2013).
14. Goetz, J. J. & Trimmer, J. M. Transcriptome sequencing of single cells with Smart-Seq. Nat Biotechnol 30(8), p. 763–5 (2012).
15. Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler transform. Nucleic Acids Res 39(2), p. 25 (2011).
16. Liu, S. J. et al. Cell-type specific features of circular RNA expression. Nature 498(7453), p. 236–40 (2013).
17. Sheng, K. et al. Effective detection of variation in single-cell transcriptomes using MATQ-seq. Nat Meth 14(3), p. 267–70 (2017).
18. Liu, S. J. et al. Single-cell analysis of long non-coding RNAs in the developing human neocortex. Genome Biol 17(1), p. 67 (2016).
19. Ma, Q. & Chang, H. Y. Single-cell profiling of IncRNAs in the developing human brain. Genome Biol 17(1), p. 68 (2016).
20. Jack, W. R. et al. Circular RNAs are abundant, conserved, and associated with ALU repeats. RNA 19(2), p. 141–57 (2013).
21. Memczak, S. et al. Circular RNAs are a large class of animal RNAs with regulatory potency. Nature 495(7441), p. 333–8 (2013).
22. Guarnero, J. et al. Oncogenic Role of Fusion-circRNAs Derived from Cancer-Associated Chromosomal Translocations. Cell 166(4), p. 1055–1056 (2016).
23. Shao, Y. & Chen, Y. Roles of Circular RNAs in Neurologic Disease. Front Mol Neurosci 9, p. 25 (2016).
24. Ghosal, S. et al. CircTracts: a comprehensive database for circular RNA potentially associated with disease and traits. Front Genet 4, p. 283 (2013).
25. Lu, W. et al. starBase v2.0: decoding miRNA-ceRNA, miRNA-ncRNA and protein-RNA interaction networks from large-scale CLIP-Seq data. Nucleic Acids Res 42(Database issue), p. D92–7 (2014).
26. Zheng, L. L. et al. deepBase v2.0: identification, expression, evolution and function of small RNAs, lncRNAs and circular RNAs from deep-sequencing data. Nucleic Acids Res 44(D1), p. D196–202 (2016).
27. Chen, X. et al. circRNADB: A comprehensive database for human circular RNAs with protein-coding annotations. Sci Rep 6, p. 34985 (2016).
28. Gao, Y., Wang, J. & Zhao, F. CIRI: an efficient and unbiased algorithm for de novo circular RNA identification. Genome Biol 16, p. 4 (2015).
29. Szabo, L. & Salzman, J. Detecting circular RNAs: bioinformatic and experimental challenges. Nat Rev Genet 17(11), p. 679–692 (2016).
30. Zhang, X. et al. Complementary sequence-mediated exon circularization. Cell 159(1), p. 134–47 (2014).
31. Song, X. et al. Circular RNA profile in gliomas revealed by identification tool UROBORUS. Nucleic Acids Res 44(9), p. e87 (2016).
32. Salzman, J. et al. Circular RNAs are the predominant transcript isoform from hundreds of human genes in diverse cell types. PLoS One 7(12), p. e30733 (2012).
33. Chen, J. et al. Single-cell SNP analyses and interpretations based on RNA-Seq data for colon cancer research. Sci Rep 6, p. 34420 (2016).
34. Salzman, J. et al. Cell-type specific features of circular RNA expression. PLoS Genet 9(9), p. e1003777 (2013).
35. Rybak, Wolf, A. et al. Circular RNAs in the Mammalian Brain Are Highly Abundant, Conserved, and Dynamically Expressed. Mol Cell 58(5), p. 870–85 (2015).
36. Subramanian, A. et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci USA 102(43), p. 15545–50 (2005).
37. Li, J. et al. Circular RNAs in cancer: novel insights into origins, properties, functions and implications. Am J Cancer Res 5(2), p. 472–80 (2015).
38. Zhang, Z., Lv, M. & Chen, J. Screening differential circular RNA expression profiles reveals the regulatory role of circTFCF25-miR-103a-3p/mir-107-CDK6 pathway in bladder carcinoma. Sci Rep 6, p. 30919 (2016).
39. Jin, X. et al. CircRNA expression pattern and circRNA-miR-mRNA network in the pathogenesis of nonalcoholic steatohepatitis. Oncotarget 7(41), p. 66455–66467 (2016).
40. Schulze, W. X., Deng, L. & Mann, P. Phosphotyrosine interactome of the ErbB-receptor kinase family. Mol Syst Biol 1, p. 0008 (2005).
41. Holbro, T. & Hynes, N. E. ErbB receptors: directing key signaling networks throughout life. Annu Rev Pharmacol Toxicol 44, p. 195–217 (2004).
42. Heuckmann, J. M., Rauh, D. & Thomas, R. K. Epidermal growth factor receptor (EGFR) signaling and covalent EGFR inhibition in lung cancer. J Clin Oncol 30(27), p. 3417–20 (2012).
43. Zhou, Q. et al. QC-Chain: fast and holistic quality control method for next-generation sequencing data. PLoS One 8(4), p. e60234 (2013).
44. Li, H. & Durbin, R. Fast and accurate read alignment with Burrows-Wheeler transform. Bioinformatics 25(14), p. 1754–60 (2009).
45. DePristo, M. A. et al. A framework for variation discovery and genotyping using next-generation DNA sequencing data. Nat Genet 43(5), p. 491–498 (2011).
46. Li, H. et al. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25(16), p. 2078–9 (2009).
47. Shannon, P. et al. Cytoscape: a software environment for integrated models of biomolecular interaction networks. Genome Res 13(11), p. 2498–504 (2003).
48. Kanemis, M. et al. KEGG: new perspectives on genomes, pathways, diseases and drugs. Nucleic Acids Res 45(D1), p. D353–D361 (2017).
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Author Contributions
K.N. designed the whole study. C.F.Z. collected datasets, C.F.Z. and S.J.Y. analyzed the data, C.F.Z., M.Z.H. explained the results, C.F.Z., S.J.Y., M.Z.H., J.H.C. and K.N. wrote the initial draft of the manuscript, all revised the manuscript.

Additional Information
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