CircleBase: an integrated resource and analysis platform for human eccDNAs

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ABSTRACT

Rapid advances in high-throughput sequencing technologies have led to the discovery of thousands of extrachromosomal circular DNAs (eccDNAs) in the human genome. Loss-of-function experiments are difficult to conduct on circular and linear chromosomes, as they usually overlap. Hence, it is challenging to interpret the molecular functions of eccDNAs. Here, we present CircleBase (http://circlebase.maolab.org), an integrated resource and analysis platform used to curate and interpret eccDNAs in multiple cell types. CircleBase identifies putative functional eccDNAs by incorporating sequencing datasets, computational predictions, and manual annotations. It classifies them into six sections including targeting genes, epigenetic regulations, regulatory elements, chromatin accessibility, chromatin interactions, and genetic variants. The eccDNA targeting and regulatory networks are displayed by informative visualization tools and then prioritized. Functional enrichment analyses revealed that the top-ranked cancer cell eccDNAs were enriched in oncogenic pathways such as the Ras and PI3K-Akt signaling pathways. In contrast, eccDNAs from healthy individuals were not significantly enriched. CircleBase provides a user-friendly interface for searching, browsing, and analyzing eccDNAs in various cell/tissue types. Thus, it is useful to screen for potential functional eccDNAs and interpret their molecular mechanisms in human cancers and other diseases.

INTRODUCTION

Extrachromosomal circular DNA (eccDNA) originates from chromosomal DNA but is independent of it. EccDNA is ubiquitous in various eukaryotes and plays multiple biological roles in different cell types (1). Researchers have known for decades that eccDNA occurs in tumor cells. However, comprehensive studies on their structure and function have returned relatively little useful information because of the limitations of available detection and analysis technology (2,3). Other researchers have exploited
recently developed high-throughput sequencing technology and eccDNA identification methods (4), revealing strong associations between eccDNA and cancers: (i) the eccDNA mediates oncogene over-expression by increasing gene copy numbers and transcript levels which lead to tumor deterioration (5–9). Tumor cells contain abundant oncogene-harboring eccDNA that is often amplified by co-amplification with adjacent enhancers. This mechanism leads to oncogene over-expression (6–8). Oncogene-harboring eccDNAs are mobile enhancers that globally amplify chromosomal transcription (5). Moreover, they have loose, open chromatin and a circular topology. Thus, multiple regulatory elements may remotely regulate transcription and enable the oncopgenes on the eccDNAs to be transcribed efficiently (9); (ii) the eccDNA mediates drug resistance in cancer cells either by amplifying the drug resistance gene on eccDNA, or by using the reversible loss of the gain-of-function mutation of the drug resistance gene carried by eccDNA (10,11); (iii) the eccDNA can drive tumor gene heterogeneity and accelerate tumor genome evolution (12). Oncogene-derived eccDNAs, which lack centromeres, may be asymmetrically delivered to progeny cells via amitosis. Cells with relatively more oncogene-derived eccDNAs have a growth advantage and are enriched in rapidly proliferating tumor cells. Computational simulation has confirmed this phenomenon (13). Neuroblastoma eccDNAs accelerate somatic genome rearrangement and oncogene remodeling via chimeric circularization and reintroduction into linear chromosomes (14).

The foregoing studies indicated that eccDNAs are promising biomarkers for cancer diagnosis and prognosis and are, therefore, research hotspots (15). Meanwhile, the promising biomarkers for cancer diagnosis and prognosis are CircleBase (http://circlebase.maolab.org) that compiles and integrates relevant databases. Additionally, 98,274,452 chromatin states were predicted using the core 25-state ChromHMM model (28). The annotation results included element annotation, targeting genes, genetic marks across all 127 reference epigenomes in the ENCODE project (29).

MATERIALS AND METHODS

eccDNA collection

CircleBase comprises 601,036 eccDNAs (candidates larger than 50M were removed) gleaned from 13 published papers on PubMed (https://pubmed.ncbi.nlm.nih.gov/) and includes the following information: (i) chromosomal localizations of eccDNAs based on reference genomes hg19 and hg38; (ii) conditions or treatments; (iii) sample types; (iv) sequencing library types and (v) validation strategies. The eccDNA localizations were collected from the Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/) (16,17) and Genome Sequence Archive (GSA, https://ngdc.cncb.ac.cn/gsa/) (18).

Gene annotation

AnnotatePeaks.pl in the HOMER package (v. 4.11.1, http://homer.salk.edu) (19) was used with its default parameters for eccDNA gene annotation based on the genomic position. The annotation results included element annotation, distance to TSS, nearest promoter ID, NCBI gene ID, nearest Refseq ID, nearest Ensembl ID, gene name, gene alias, gene description and gene type.

Targeting genes

3,049,674 enhancer-target pairs from human variation annotation database (VARAdb) (20) were collected to determine the target genes of the eccDNAs located in the enhancers and super-enhancers. There were 9,879,737 enhancer-target networks in 935 samples determined using the JEME algorithm (https://github.com/yiplabcuhk/JEME) (21) and 284,834 enhancer-gene links identified using the GeneHancer database (22).

Regulatory elements

Regulatory elements are anchored by chromatin looping to gene promoter regions to regulate gene transcription. Enhancers and super-enhancers are positive regulatory elements that maintain cell type-specific gene expression and control cell fate during development (23). There were 65,950 super-enhancers from dbSUPER (24), 2,839,656 enhancers from EnhancerAtlas (25), 22,601,146 enhancers/super-enhancers from SEA (26) and 331,146 super-enhancers from SEdb (27). They were collected in CircleBase, and their target genes were annotated according to their source databases. Additionally, 98,274,452 chromatin states were predicted using the core 25-state ChromHMM model (28). ChromHMM was trained using the data imputed for 12 epigenetic marks across all 127 reference epigenomes in the ENCODE project (29).

Chromatin interactions

Chromatin loops mediated by the CTCF/cohesin complex connect regulatory elements to their target genes. In CircleBase, 28,442,796 chromatin interaction records were collected from three databases: (i) 25,222,085 pairs of chromatin interactions were identified using the EpiTensor algorithm on Oncobase (30). These included 2,847,794, 5,691,699 and 16,682,592 promoter–promoter, enhancer–promoter and enhancer–enhancer interactions, respectively. (ii) 3,095,881 pairs of chromatin interactions were collected from the 4DGenome database (31). They were experimentally determined via Hi-C, ChIA-PET, IM-PET, 3C, 4C and 5C. (iii)
124,830 ChIA-PET narrow peaks were detected from ENCODE (29) in MCF-7, HeLa-S3, K562, NB4, and HCT116 cancer cells.

Logistic function for score normalization

The scores were differently defined in the various annotation databases. Therefore, they were transformed to harmonize their ranges within [0,1]. The Z-score for each record in each dataset was calculated as follows:

$$Z(x) = \frac{x - \bar{x}}{SD(x)}$$

where \(\bar{x}\) and \(SD(x)\) are the mean and standard deviation for all samples, respectively. A logistic function was then used to convert the Z-score as follows:

$$f(Z) = \frac{1}{1 + e^{-z}}$$

where \(Z\) is the Z-score of each sample, \(e\) is the base of natural logarithm.

Diffusion system for prioritizing target genes

The network-based diffusion algorithm PageRank was used to prioritize the target genes for each eccDNA (30). An improved version of PageRank was applied as follows:

$$P R(u) = (1 - d) + d \times \sum_{v \in B(u)} P R(v) \times weight_{v \rightarrow u}$$

where \(Weight_{v \rightarrow u}\) is used to measure the weight of the edge. \(Weight = 1\) for the edge derived from the enhancer/super-enhancer interactions determined by the regulatory elements section. \(Weight = 0.5\) for the edge derived from the chromatin connections determined by the chromatin interactions section. In the targeting genes section, the original enhancer-target scores derived from VARAdb and JEME were used as the weight values, and the enhancer-target scores from GeneHancer were normalized to [0, 1] via the foregoing logistic function. The mean weight scores were used as ultimate edge scores when a single eccDNA was implicated in >1 connection record.

Regulatory networks

Regulatory networks were constructed by combining the enhancer-gene pairs from the targeting genes, regulatory elements, and chromatin interactions sections. Regulatory element targets overlapping with eccDNA were defined as eccDNA targets. Redundant eccDNA–gene pairs were removed. Genes interacting with eccDNA were ranked using Google PageRank and circle sizes were positively correlated with PageRank scores (32). An interactive view was prepared for the gene prioritization related to certain eccDNAs in a two-layer network ranked by PageRank score in the regulatory network section (30). PageRank ranking was applied to all genes regulated by certain eccDNA and all other eccDNAs regulating them. The network view showed the two-layer networks. The eccDNAs or genes related to the sub-networks were listed in the table following the interactive view.

Functional enrichment

A functional enrichment analysis was performed using the R package clusterProfiler version 4.0.2 (33). A gene ontology analysis was performed with the clusterProfiler enrichGO function (34). Kyoto Encyclopedia of Genes and Genomes pathways were analyzed with the clusterProfiler enrichKEGG function (35). Terms were considered significantly enriched when their Benjamini–Hochberg-adjusted \(P\)-values were <0.05 (36).
Chromatin accessibility

Chromatin accessibility reflected both aggregate TF binding and the regulatory potential of a genetic locus (37). In CircleBase, (i) 52 078 883 accessible regions were detected by ATAC-seq in over 1400 samples from ATACdb (38); (ii) 3 181 274 accessible regions were detected by ATAC-seq in Cistrome (39); (iii) 1 051 532 accessible regions were detected by ATAC-seq in 23 cancer types from The Cancer Genome Atlas (TCGA) (40) and (iv) 62 154 007 accessible regions were detected by DNase-seq in 243 samples from ENCODE (29).

Epigenetic regulations

There were 156 379 641 peaks collected from 7734 ChIP-seq samples of 952 TFs in ChIP-Atlas (41), Cistrome (39), ENCODE (29), GTRD (42) and ReMap (43). Moreover, 5 911 338 and 64 669 729 histone modification peaks were obtained from 185 and 979 ChIP-seq samples from ENCODE (29) and Roadmap (44), respectively. The targets of ChIP-seq experiments from ENCODE (29) included H2A.Z, H3K27ac, H3K27me3, H3K36me3, H3K4me1, H3K4me2, H3K4me3, H3K79me2, H3K9ac, H3K9me1, H3K9me3, H4K20me1, and H4K12ac. The targets of ChIP-seq experiments from Roadmap included H2A, H2AK5ac, H2AK9ac, H2BK120ac, H2BK12ac, H2BK15ac, H2BK20ac, H2BK5ac, H3K14ac, H3K18ac, H3K23ac, H3K27me2, H3K27ac, H3K27me3, H3K36me3, H3K4ac, H3K4me1, H3K4me2, H3K4me3, H3K56ac, H3K79me1, H3K79me2, H3K9ac, H3K9me1, H3K9me3, H3T11ph, H4K12ac, H4K20me1, H4K5ac, H4K8ac and H4K91ac.

Genetic variants

Genetic variants were collected from 1 245 959 177 human single-nucleotide variants (SNVs) in dbSNP release 155 (45), 8 775 948 common single-nucleotide polymorphisms (SNPs) from the Genome Aggregation Database (gnomAD) Project v. 2.1 (46), 27 260 726 SNPs from the genome-wide association study (GWAS) catalog (47) and 314 238 risk SNPs from GWASdb v. 2.0 (48). Disease-related genetic variants including 81 666 495 somatic mutations of human cancers from OncoBase (30) and 670 082 de novo mutations from the Gene4Denovo database (49), were also collected.

Expression quantitative trait locus (eQTL)

The eQTL are genomic loci that regulate gene expression levels and play vital roles in deciphering gene regulation and spatiotemporal specificity (50). Correlation between genotype and tissue-specific gene expression level may be used to interpret the effect of gene variant on gene expression in human tissues or cancers. Here, 71 478 479 significant SNP–gene pairs (false discovery rate (FDR) < 0.05) in 49 human tissues were compiled from the GTEX Project v. 8.0 (51). Additionally, 1 412 029 significant cis-eQTL-gene and trans-eQTL-gene pairs in 33 cancer types were collected from the PancanQTL database (52).

Interactive circular visualization

High-resolution chromatin interactions, TF binding clusters, somatic mutations, enhancers, super-enhancers, and their predicted targets were illustrated in a circular ideogram layout plotted with BioCircos (http://bioinfo.ipb.ac.cn/biocircos/index.php) (53). The ideogram was then implemented for circular visualization of various biological data such as genomic features, genetic variants, gene expression, and biomolecular interactions.

ecDNA prioritization scoring system

The number of annotated hits (records) assigning a score to each corresponding ecDNA was used for each regulatory category. Considering that ecDNA has $k$ hits per regulatory category ($F$), and $\mu$ and $\sigma$ are the fitted parameters of the corresponding Gaussian model, the ecDNA score in this category was calculated as follows:

$$Score_F = -\log_{10}\left(\int_{k}^{\infty} \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}} dx\right)$$

(4)

The score of each category was normalized to [0, 1] using the foregoing logistic function. The final score $S$ of each ecDNA would be the average of normalized scores for all six regulatory categories (Equation 5) since normalized scores between some categories are not independent (Supplemental Figure S1). The calculation of the scoring system was implemented using the SciPy package v1.6.3 in Python v3.7.6 (54).

$$S = \frac{1}{6} \sum_{F=1}^{6} Score_F$$

(5)

System design and database construction

CircleBase (http://circlebase.maolab.org) was developed by combining jQuery with the PHP-based web framework CodeIgniter (https://codeigniter.com). It was based on previously reported databases and platforms (30,50,55,56). All datasets in CircleBase were stored either in MySQL database or as flat files. Academic users may freely access related data and analytical results through the web interface. The LiftOver module in the UCSC toolkit (https://genome.ucsc.edu/cgi-bin/hgLiftOver) (57) was used to convert genomic coordinates among various versions of the human genome. In details, ecDNA locations from the paper PMID 34193237 were lifted over from hg38 to hg19 while the rest were lifted over from hg19 to hg38. In addition, super-enhancer locations from the SEA database were lifted over from hg38 to hg19.

Data and code availability

The accession, DOI, version and permalink information of the foregoing databases was listed in Supplementary Table S1. To make the codes open-access, we deposited the source codes of CircleBase in GitHub (https://github.com/leishenggit/CircleBase). The full tables of human ecDNAs are available in the Download module of CircleBase (http://circlebase.maolab.org/welcome/download).
Figure 2. Percentage of eccDNAs in each category and distribution of ranking scores. (A) Percentage of eccDNAs in three cell groups including cancer cell line, cancer tissue, and healthy individual. (B) Percentage of eccDNAs in different library types such as WGS, Circle-Seq, ATAC-seq, WES and ChIA-PET. (C) Percentage of eccDNAs in different cell lines such as ES2, LnCap, U937, OVCAR8, C4-2, HeLaS3, PC-3 and C4-2B. (D) Percentage of eccDNAs in different human tissues such as muscle, plasma, connective tissue, brain, breast, esophagus and lung. (E) Density distribution of eccDNAs in different ranking scores.
Figure 3. Web interface of the Search module in CircleBase.

Figure 4. Functional enrichment analysis of the top 1000 eccDNAs. (A) Functional enrichment analysis of 791 eccDNAs from cancer cells. (B) Functional enrichment analysis of 209 eccDNAs from normal tissues.
Figure 5. Illustration of the eccDNA ecc595499 in gastric cancer. (A) BioCircos view of targeting genes. (B) BioCircos view of chromatin interactions. (C) Regulatory network. The interacting genes with the eccDNA are ranked by Google PageRank and the size of the circle is positively correlated with the PageRank score. (D) Molecular function enrichment of eccDNA ecc595499 targeting genes.

DATABASE FEATURES AND APPLICATIONS

Statistics and features

CircleBase compiled available eccDNAs \((n = 601 \, 036)\) from public resources and comprehensively interpreted those in various cancer cell lines \((n = 423 \, 018)\), cancer tissues \((n = 6498)\), and healthy individuals \((n = 171 \, 596)\) (Figure 2A). These eccDNAs were detected by whole genome sequencing (WGS, \(n = 427 \, 064\)), Circle-Seq \((n = 145 \, 588)\), cf-eccDNA-seq \((n = 25 \, 995)\), ATAC-seq \((n = 1783)\), whole exome sequencing (WES, \(n = 467)\) and others (Figure 2B). We collected 114 611 eccDNAs from ES2, 84 629 from LnCap, 69 420 from U937, 57 780 from OVCAR8, 41 221 from C42 and 38 876 from HeLaS3 cell lines (Figure 2C). We collected 136 619 eccDNAs from the muscle, 34 977 from the plasma, 34 977 from the plasma, 939 from the connective tissue, 754 from the brain, 712 from the breast, 648 from the esophagus, 599 from the lung, 435 from the skin, 427 from the stomach, 362 from the bladder, 301 from the head and neck, 340 from the ovaries, 250 from the uterine corpus endometrial, 153 from the liver, 157 from the pancreas, 149 from the prostate, and others (Figure 2D).

Unlike our previous study (55), the hits numbers of eccDNAs per chromosome were not suitable for the Poisson distribution (Supplemental Figures S2-S7). The parameter lambda of the Poisson distribution is approximated by the average value of the observed data, and then it is substituted into the R function \(rpois\) to generate the theoretical values. But the hits numbers per chromosome could be fitted to a Gaussian distribution by using Box–Cox transformation (Supplemental Figures S8-S13). The parameters of the normal distribution, that is, the mean and variance are approximated by the mean and variance of the observed data, and then they are substituted into the R function \(rnorm\) to generate the theoretical values. Importantly, the mean values of six scores after logistic transformation conform to a normal distribution with range of \([0, 1]\) (Figure 2E). Therefore, CircleBase helps prioritize putative functional eccDNAs by incorporating high-throughput experimental datasets from ENCODE (29), computational predictions, and manual annotations. It classifies these genetic and epigenetic annotations into targeting genes, epigenetic regulations, regulatory elements, chromatin accessibility, chromatin interactions and genetic variants (Figure 1). It predicts regulatory networks between eccDNAs and genetic/epigenomic fac-
Figure 6. Illustration of the eccDNA ecc597923 in ovarian cancer. (A) BioCircos view of targeting genes. (B) Regulatory network. The interacting genes with the eccDNA are ranked by Google PageRank and the size of the circle is positively correlated with the PageRank score. (C) BioCircos view of chromatin interactions. (D) Biological process enrichment of eccDNA targeting genes.

tors by integrating relevant databases. These data sources are combined to score eccDNAs and help interpret the potential functions of the eccDNAs in the human genome.

Website interface

CircleBase comprises the Home, Search, Stats, Manual, Download, Links and Contact modules. A brief introduction of CircleBase and the workflow are shown in the Home module. Data are integrated in the Search module of CircleBase and the workflow are shown in the Home module. Data are integrated in the Search module of CircleBase and can be searched by gene symbol, genomic region, eccDNA ID, NCBI gene ID and Ensembl gene ID. Moreover, examples are provided for new users. When users search their target eccDNAs, the target genes, regulatory elements, chromatin interactions, regulatory networks, functional enrichments, chromatin access, epigenetic regulations, genetic variants and eQTL, and the summary are displayed at the click of a mouse (Figure 3). In the Stats module, eccDNA distributions under various criteria including cell line and cell group, tissue, and library type are exhibited in a pie chart. In the Manual module, steps of how to use CircleBase are shown. In the Download module, all eccDNA data used are available for user download. In the Links Module, related databases such as TCGA (40), International Cancer Genome Consortium (ICGC) (58), and others are provided. In the Contact module, the e-mail addresses and research fields of related authors are described in detail, enabling users to communicate with our team.

Significance and applications

The top 1000 eccDNAs scored by the ranking system were used to perform functional enrichment analyses (59) including 791 and 209 eccDNAs from cancer cells and normal tissues, respectively. The top-ranked cancer cell eccDNAs were enriched in oncogenic pathways (Figure 4A) including Rap1 signaling (hsa04015), Ras signaling (hsa04014), Melanoma (hsa05218), MAPK signaling (hsa04010), PI3K-Akt signaling (hsa04151), Regulation of actin cytoskeleton (hsa04810), Phospholipase D signaling (hsa04072), Gastric cancer (hsa05226), Pathways in cancer (hsa05200) and Circadian entrainment (hsa04713). By con-
trast, eccDNAs from normal tissues had no significant enrichment (Figure 4B). The foregoing results validated the ranking system used in CircleBase.

Here, we considered eccDNA ecc595499 (chr10:81044184–81134986) in gastric cancer and eccDNA ecc597923 (chr2:176959007–177342179) in ovarian cancer as examples (60). CircleBase showed the targeting gengs, chromatin interactions in their loci and flanking regions, and regulatory network (Figures 5A–C; Figures 6A–C). Our platform predicted several targeting genes of ecc595499 enriched in multiple molecular functions including carbohydrate binding, S100 protein binding, MHC class II protein complex binding, translation elongation factor activity, cyclosporin A binding, MHC protein complex binding, peptidyl-prolyl cis-trans isomerase activity, cis-trans isomerase activity, calcium-dependent phospholipid binding, nuclear receptor coactivator activity, ribosome binding, and drug binding (Figure 5D). Our platform disclosed dozens of ecc597923 targeting genes enriched in multiple biological processes related to embryonic organ development such as appendage morphogenesis, limb morphogenesis, appendage development, limb development, embryonic limb morphogenesis, and embryonic appendage morphogenesis (Figure 6D). Experimental validation of eccDNA function is currently in progress and we believe that by using the regulatory networks and functional predictions from CircleBase, researchers will be able to elucidate the mechanisms underlying the functions of eccDNAs in human cancers and other diseases in the near future.

DISCUSSION AND PERSPECTIVES

Cancer cells can rapidly adapt to changes in the tumor micro-environment by amplifying the oncogenes on their eccDNA (61). Numerous experimental methods have been developed to detect eccDNA including Circle-Seq based on next-generation sequencing (62) and SMOOTH-seq based on third-generation sequencing platforms (63). Several computational methods have been developed to analyze eccDNAs including AmpiclonArchitect (64), ECdetect (6) to identify eccDNAs from whole-genome sequencing data, and Circle-Map (65) and Circle_finder (66) to detect eccDNAs in Circle-seq and ATAC-seq data, respectively. Integrated databases and annotation platforms are required for human eccDNAs. However, none of these have been developed or published to date. As far as we know, CircleBase is the first database for eccDNA and has several advantages. It (i) is fitted with a highly interactive visualization function for eccDNAs and their related annotations; (ii) has a ranking system based on a Gaussian distribution model for better decision-making and (iii) provides comprehensive eccDNA annotations as follows:

i) It furnishes annotations for eccDNA targeting genes based on bioinformatics predictions by JEME (21) and EpiTensor in the OncoBase platform (30).

ii) It incorporates epigenome information from ENCODE (29) and the Roadmap (44) epigenomics project with the chromatin status of the eccDNA locus in the linear genome.

iii) It fine-maps the genetic basis of the eccDNA using disease-related and common SNPs from TCGA (40), gnomAD (46), dbSNP (45), GWASdb (48) and others.

iv) It annotates eccDNAs to regulatory elements, such as promoters, enhancers, and super enhancers via ChromHMM (28), EnhancerAtlas (25), dbSUPER (24) and others.

v) It measures the chromatin accessibility of the eccDNA locus with ATACdb (38), Cistrome (39) and others.

vi) It establishes chromatin interaction networks for the eccDNA locus using Hi-C processed data from 4DGenome (31), OncoBase (30) and others.

In conclusion, CircleBase curates eccDNAs in the human genome from various cell types and provides a scoring system to prioritize the eccDNAs based on comprehensive annotations. Our aims are to extend the ranges of annotations and cell types and continue updating CircleBase.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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