Identification of a lncRNA-associated competing endogenous RNA-regulated network in clear cell renal cell carcinoma

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Abstract. Long noncoding RNAs (lncRNAs) act as competing endogenous RNAs (ceRNAs) in the regulation of gene expression in various physiological and pathological processes. The present study aimed to explore the lncRNA-miRNA-mRNA interactions in clear cell renal cell carcinoma (ccRCC) using comprehensive bioinformatics analysis. RNA-seq data were downloaded from the TCGA Data Portal, and the differentially expressed lncRNAs (DELncRs), miRNAs (DEMriRs), and mRNAs (DEGs) between tumoral and control samples were identified using the edgeR package. The correlations between the DEMiR/DELncR expression levels and clinical features were evaluated using nonparametric regression analysis. The Kaplan-Meier method was used to identify DELncRs associated with overall survival time. Then, the DELncR-DEMriR interaction pairs were predicted using mirCode and the starBase v2.0 database, and DEMiR-DEG pairs were predicted using the miRTarBase database. Then, a ceRNA-regulated network of ccRCC was constructed based on these interactions. Genes in the network were also assigned to functional categories in the KEGG pathway database. A total of 1,573 DEGs, 37 DELncRs, and 62 DEMiRs were identified. Moreover, several DELncRs were significantly associated with patient clinical variables; for example, TCL6 was significantly associated with tumor grade and AJCC pathological stage. Next, 38 pairs of DELncR-DEMriR interactions (13 DELncRs and 8 DEMiRs) were identified. Among the 8 DEMiRs that target DELncRs, six were found to target DEGs. Based on the identified DELncR-DEMriR interactions and DEMiR-DEG interactions, a ceRNA-regulated network comprising 203 nodes and 221 edges was constructed (with MIC >0.15 and MIC-p2 >0.15). The novel lncRNAs, DGCR5, MYCNOS, and PART1 may participate in the progression of ccRCC through cytochrome P450-mediated drug metabolism.

Introduction

Renal cell carcinoma (RCC) is one of the 10 most common cancers, accounting for 2-3% of all adult malignancies (1). Clear cell renal cell carcinoma (ccRCC) is the most common subtype of RCC, and approximately 75-80% of patients with RCC are diagnosed with ccRCC (2). Current therapies for renal cancer include conventional chemotherapy and radiation, however, the prognosis is poor (3). Therefore, to aid in the development of effective treatments for ccRCC, it is necessary to investigate the underlying molecular mechanism involved in the pathogenesis of ccRCC.

Only 2% of RNAs encode proteins in human cells. Many transcripts that do not encode proteins have recently been identified in large-scale genomic studies (4,5). Although these RNAs are not translated, they play critical roles in regulating transcriptional and non-transcriptional processes (6). These non-coding RNAs (ncRNAs) can be broadly classified as small ncRNAs (<200 nucleotides, including microRNAs (miRNAs), siRNAs, and piRNAs) or long non-coding (lncRNAs) (>200 nucleotides). miRNAs are the most widely studied subclass of small ncRNAs, and they can post-transcriptionally regulate the expression of multiple genes via the imperfect complementarity to their target mRNA transcripts (7). lncRNAs have been revealed to act as competing endogenous RNAs (ceRNAs), where the lncRNAs bind to and keep miRNAs from their cognate mRNA targets (11,12). Hence, the miRNAs can be prevented from mediating miRNA-mediated repression (11,12). A previous study indicated that small concentration changes in miRNA-mRNA or miRNA-ceRNA

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pairs can substantially affect the regulation of the gene expression network (13). For example, the lncRNA HULC has been previously shown to act as an endogenous sponge and inhibit miR-372, which plays an important role in tumorigenesis in liver cancer (14). The muscle-specific lncRNA, linc-MDI, regulates muscle differentiation by sequestering miR-133 and miR-135 and thus modulating the expression of MAMl1 and MEF2C (15). Recent advances in the study of the functions and mechanisms of lncRNAs in physiological and pathological processes have led to expanding the possible manner in which lncRNAs function within the already complex system of miRNA-mediated gene regulation.

Although the expression profile of mRNA and lncRNA transcripts in ccRCC has been explored and several lncRNAs, such as HIF-1alpha-AS1, H19, KCQ1OT1, and MALAT1, have been reported to be involved in renal cancer (16-18), the regulation mechanism of lncRNA-miRNA-mRNA in ccRCC has not been elucidated. In this study, we downloaded the data for an important cohort encompassing more than 300 samples from the Cancer Genomes Atlas (TCGA) and used a bioinformatics method to predict the regulation mechanism of lncRNAs-miRNAs-mRNAs in ccRCC.

Materials and methods

Data

RNA-seq data collection and processing. The TCGA database is a comprehensive, publicly available, and vast source of information on cancer genetic and epigenetic profiles. The TCGA RNA-seq expression datasets (level 3) for ccRCC were downloaded from the TCGA Data Portal (https://tcga-data.nci.nih.gov/tcga/) (June 3, 2016), which contained 606 mRNA expression profiles and 326 miRNA expression profiles. Since TCGA level 3 data typically represents aggregated, normalized, and/or segmented data, the level 3 data downloaded was analyzed without normalization. Based on the barcodes, the RNA-seq samples were matched to miRNAs, and a total of 326 samples, including 254 ccRCC samples and 71 normal controls, were obtained.

lncRNA and mRNA sequencing was performed on the Illumina Hiseq 2000 platform, and miRNAs were sequenced on the RNASeqV2 platform (https://www.cancer.gov/about-nci/organization/ccg/research/structural-genomics/tcga/using-tcga/technology/Illumina-HiSeq2000-Data-Sheet). RNA-seq reads were processed using RNA-Seq by Expectation-Maximization (RSEM). In addition, lncRNAs and mRNAs were annotated and identified on the website of the HUGO gene nomenclature committee (HGNC, http://www.genenames.org/), which contains annotation information on 2,775 lncRNAs and 19,004 protein-coding genes.

Analysis of differentially expressed mRNAs (DEGs), IncRNAs (DEIncRs), and miRNAs (DEmiRs). The differential expression of mRNAs, IncRNAs, and miRNAs was analyzed using the edgeR package (19), which is an R Biocconductur package that provides statistical routines for examining the differential expression of replicated count data in RNA-seq data. This package is based on an overdispersed Poisson model (20) to account for the variability at both the biological and technical levels. In addition, the Empirical Bayes methods (21) in this package were used to moderate the overdispersion across transcripts, improving the reliability of inferences. The nominal P-values were estimated based on the method of Benjamini and Hochberg (22), using the multtest package (http://www.bioconductor.org/packages/release/bioc/html/multtest.html) in R. Only RNAs with a false-discovery rate (FDR) <0.05 and logFC (fold change)>1 were selected as DEGs, DEIncRs, or DEmiRs.

Correlation analysis between the clinical characteristics and expression of DemiRs and DEIncRs. The clinical characteristics of patients with ccRCC were extracted from the downloaded data, including sex (female vs. male), age at diagnosis (≥61 years vs. <61 years), tumor grade (G3/G4 vs. G1/G2), tumor status (tumor-burden state vs. tumor-free state), American Joint Committee on Cancer pathologic stage (III/IV vs. I/II), tumor metastasis (yes vs. no), and overall survival time. Then, the DEIncRs/DemiRs that were significantly correlated with these clinical characteristics were identified using nonparametric regression analysis. A survival curve was constructed using the Kaplan-Meier method (23), and comparisons using the log rank test were performed between patients with upregulated DEIncRs and those with downregulated DEIncRs.

Construction of a ceRNA-regulated network. The lncRNA targets of human miRNAs were downloaded from the mirCode database (http://www.mircode.org/), a comprehensive searchable map of putative miRNA target sites across the complete GENCODE annotated transcriptome established by Jeggari et al. (24), and starBase v2.0 (http://starbase.sysu.edu.cn/), a platform comprised of 111 CLIP-Seq data sets and focusing on protein-lncRNA, protein-snRNA, protein-mRNA and protein-pseudogene interactions. The miRNA targets from the miRtarBase (http://mirtarbase.mbc.nctu.edu.tw), which is the most updated collection of miRNA-target interactions, were identified.

According to the identified miRNA-lncRNA interactions and miRNA-mRNA interactions, a comprehensive lncRNA-miRNA-mRNA network was constructed. Then, the interactions in the lncRNA-miRNA-mRNA network were further identified by calculating maximal information coefficients (MICs) with cut-off values of MIC >0.15 and MIC-p >0.15.

Functional enrichment of DEGs in the ceRNA-regulated network. The DEGs in the ceRNA-regulated network were assigned to functional categories in the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway database (25) (https://www.genome.jp/kegg/), and the pathways significantly associated with ccRcc were obtained. Enrichment P-values were calculated using Fisher's exact tests using the equation below. Then, the functional pathways in the ceRNA-regulated network in which the DEGs were enriched were identified.

\[
p = 1 - \sum_{i=0}^{[M]} \binom{M}{i} \binom{N-i}{K-i} \binom{N}{K}
\]

Here, N represents the number of genes in the whole genome; M represents the number of genes enriched in KEGG
pathways; and K represents the number of differentially expressed genes. The Fisher’s score represents the probability that at least x genes of K differentially expressed genes are enriched in KEGG pathways.

Results

Identification of DEGs, DelncRs, and DEmiRs. IncRNAs and mRNAs were annotated and identified on the HUGO gene nomenclature committee website. A total of 819 IncRNAs and 18,137 protein-coding transcripts were identified. Subsequently, using an FDR statistic <0.05 and IlogFC; (fold change)>1, the differential expression analysis identified 1,573 DEGs and 37 DELncRs in the RNA-seq data and 62 DEmiRs in the miRNA-seq data. A heatmap representing the expression of the identified DELncRs, DEGs, and DEmiRs is presented in Fig. 1. The ccRCC samples were obviously separated from the control samples, suggesting that the DEGs represent a large difference between the tumor and control samples.

Identification of DemiRs and DelncRs that are significantly correlated with clinical characteristics. The correlation analysis between the clinical characteristics of patients and DemiRs indicated that the expression level of hsa-miR-216b was associated with two clinical variables, sex and tumor grade (Table I).

Moreover, the correlation between the clinical characteristics of patients and the expression of identified DelncRs was explored. The results are presented in Table II. The expression of DGCR9 and TTTY4C was significantly

Figure 1. Heatmap representing the expression of differentially expressed (A) IncRNAs, (B) mRNAs, and (C) miRNAs. IncRNAs, long noncoding RNAs; miRNAs, microRNAs.
correlated with three clinical variables, sex, tumor status, and metastasis. TCL6 was revealed to be significantly associated with tumor grade and AJCC pathological stage. In addition, tumor metastasis was related to the upregulated expression of BPES1C1, C20orf203, C7orf71, C2orf48, DGCR9, DGCR5, and DKFZp434J0226 and the downregulated expression of MYCNOS, PP14571, TTTY4C, and PART1. These results revealed certain candidate lncRNAs for ccRCC, such as C7orf71, C2orf48, DGCR9, DGCR5, and TTTY4C, and also suggested a sex difference in the stage distribution of ccRCC.

Kaplan Meier analysis was conducted on all the 37 DElncRs that were significantly associated with patient clinical characteristics. As a result, 7 DElncRs that were significantly associated with overall survival time of patients with ccRCC were identified, TTTY4C, PSORSIC3, C2orf48, HPYR1, PWRN1, C15orf56, and PART1. All of these DElncRs were negatively correlated with patient overall survival, as patients with lower expression of these DElncRs had longer survival times (Fig. 2).

**Identification of DElncR-DEmiR interaction pairs.** Predictions using miRcode and starBase revealed a total of 353 pairs of DElncR-miRNA interactions and 38 pairs of DElncR-DEmiR interactions. An interaction network of 13 DElncRs and 8 DEmiRs was constructed (Fig. 3). miR-122, miR-129, and miR-216b targeted more lncRNAs than the other miRNAs. miR-122 had a relationship with 8 DElncRNAs, including MYCNOS, PART1, and PWRN1. miR-129 targeted 7 DElncRNAs, including DGCR5, HPYR1, MYCNOS, PART1, and PWRN1. miR-216b had 7 targets, BPES1C1, DGCR5, FAM138E, HPYR1, PVT1, PWRN1, and SFTA1P.

**Identification of DEmiR-DEG interaction pairs.** As aforementioned, a total of 8 DEmiRs were revealed to be DElncR-targeting miRNAs. Then, the DEmiR-mRNA
Figure 2. Kaplan-Meier survival curves for 7 identified differentially expressed lncRNAs, including (A) C15orf56, (B) HPYR1, (C) C2orf48, (D) PSORS1C3, (E) TTTY4C, (F) PART1, and (G) PWRN1. lncRNAs, long noncoding RNAs.
Figure 3. Interaction network of identified DElncRs and DEmiRs. The white diamonds represent differentially expressed lncRNAs; the blue hexagons represent differentially expressed miRNAs. The full lines represent the lncRNA-miRNA interactions. DElncRs, differentially expressed lncRNAs; DEmiRs, differentially expressed miRNAs; miRNAs, microRNAs. The red hexagon indicates a tumor-associated gene included in the allOnco database.

Figure 4. Interaction network of identified DEmiRs and DEGs. (A) miR-122-DEG interaction network; (B) miR-138-DEG interaction network; (C) miR-141-DEG interaction network; (D) miR-184-DEG interaction network; (E) miR-206-DEG interaction network; and (F) miR-216b-DEG interaction network. The blue hexagons represent differentially expressed miRNAs; the white circles represent differentially expressed genes; the red hexagons represent cancer-related genes from the allOnco database. The full lines represent the mRNA-miRNA interactions. DEmiRs, differentially expressed miRNAs; DEGs, differentially expressed genes; miRNAs, microRNAs.
interactions were collected from the miRTarBase database, and 6 out of 8 DEMiRs were revealed to target DEGs: hsa-miR-122 targeted 26 DEGs, hsa-miR-216b targeted 30 DEGs, hsa-miR-141 targeted 54 DEGs, hsa-miR-206 targeted 15 DEGs, hsa-miR-138 targeted 25 DEGs, and hsa-miR-184 targeted 32 DEGs (Table III). Collectively, these data indicated that hsa-miR-122 and hsa-miR-216b may be important in ccRCC. In addition, hsa-miR-141 had the most targets, followed by hsa-miR-184, also indicating their crucial role in ccRCC.

Among the 182 DEMiR targets, 130 DEGs were identified as cancer-related genes after comparison with the all onco data-base (http://www.bushmanlab.org/links/genelists), a collection of 2,125 cancer-related genes. The DEMiR-DEG interaction network is presented in Fig. 4.

Construction and functional annotation of a ceRNA-regulated network. A comprehensive lncRNA-miRNA-mRNA network was established based on the identified DEMiR-DEG interactions and DEMiR-DEG interactions. By calculating MICs, with MIC >0.15 and MIC-p^2 >0.15, a ceRNA-regulated network comprised of 203 nodes and 221 edges was constructed (Fig. 5). This network indicated the roles of hsa-miR-141, hsa-miR-184, hsa-miR-122, hsa-miR-216b, hsa-miR-138, and hsa-miR-206 in the ceRNA-regulated network in ccRCC.

To study the function of DEGs in the ceRNA-regulated network, KEGG pathway and gene ontology (GO) catalogue annotation were performed. These DEGs were enriched in 8 KEGG pathways (Fig. 6A), such as metabolism of xenobiotics by cytochrome P450, retinol metabolism, and drug metabolism, and 17 GO terms, such as intrinsic to membrane, extracellular region, extracellular region part, transcription factor activity, ion transport, and proteolysis (Fig. 6B).

Discussion

Human IncRNAs have been revealed to act as competitors of mRNAs for miRNA binding in RNA-RNA cross-talk networks and regulate mRNA expression at the post-transcriptional level. Although certain differentially expressed IncRNAs and mRNAs have been identified in ccRCC by expression profiling and a branch of the MALAT1/miR-200s/ZEB2 pathway, and the functional triplets consisting of IncRNAs, mRNAs, and miRNAs, have been proposed to regulate the progression of ccRCC, a systematic analysis of the ceRNA-regulated network in ccRCC has not been performed. To understand the roles of the ceRNA-regulated network in the pathology of ccRCC, a transcriptome-wide identification and characterization of differentially expressed RNAs, including IncRNAs, mRNAs, and miRNAs was performed, and a comprehensive IncRNA-miRNA-mRNA network was constructed based on identified IncRNA-miRNA interactions and miRNA-mRNA interactions downloaded from an online database.
To the best of our knowledge, the present study represents the first comprehensive description of the ceRNA-regulated network in ccRCC, as identified through the bioinformatics data portal. The present study provides new insights that will aid in understanding the molecular mechanism involved in ccRCC.
analysis of RNAseq data in a large patient cohort that encompasses 254 ccRCC tumor samples and 71 normal renal tissue samples. Several novel findings were obtained from our in-depth genomic analysis.

First, the expression of certain differentially expressed lncRNAs, such as C2orf48, DGC5, DGCR5, DGF2p4, D4J2, MYCNOS, PPI4571, TTTY4C, and PART1, was correlated with tumor metastasis; moreover, the expression of TTTY4C, PSORS1C3, C2orf48, HYPR1, PWRN1, C15orf56, and PART1 was negatively correlated with patient overall survival. The expression level of miR-216b was revealed to be associated with sex and tumor grade in patients with ccRCC (26). Notably, TTTY4C and certain other lncRNAs were also revealed to be related to sex, which is consistent with the sex-dependent manner of ccRCC (27). All these identified lncRNAs have not been previously reported to be associated with ccRCC. DGCR5 has been revealed to be involved in the regulation of the proliferation, migration, and invasion of lung cancer (28) and lung adenocarcinoma (29). MYCNOS has been reported to cooperate with a crucial transcription factor, CTCF-binding factor, to promote neuroblastoma progression through MYCN expression (30). PSORS1C3 has been demonstrated to be expressed in pluripotent and tumor cell lines (31). Therefore, further studies are required to clarify the role of these identified lncRNAs in ccRCC.

Second, a comprehensive lncRNA-miRNA-mRNA network in ccRCC was constructed, which revealed the roles of hsa-miR-141, hsa-miR-184, hsa-miR-122, hsa-miR-216b, hsa-miR-138, and hsa-miR-206 in the ceRNA-regulated network in ccRCC. The identified miRNAs in the ceRNA-regulated network, including miR-141 (32,33), miR-138 (32-34), miR-206c (35), miR-122 (36), miR-184 (37), and miR-216b (26), have been reported to play important roles in tumor progression in human ccRCC. In addition to these miRNAs, lncRNAs were also identified in the ceRNA-regulated network. As aforementioned, DGCR5 plays an important role in lung cancer (28,29) and was revealed to target miR-216b, miR-129, miR-206, miR-141, and miR-138 in the present study. MYCNOS was revealed to target miR-129, miR-122, and miR-138. PART1 has also been revealed to target miR-129, miR-206, miR-219, miR-206, miR-141, and miR-138. Based on our results, the ceRNA-regulated network revealed how these competing modules (lncRNA-miRNA-mRNA) play a role in the pathogenesis of ccRCC and indicated the roles played by lncRNAs in the disease.

Moreover, the transcripts in the ceRNA network of ccRCC were revealed to be enriched in several pathways such as metabolism of xenobiotics by cytochrome P450, retinol metabolism, and drug metabolism and critical biological processes such as intrinsic to membrane, extracellular region, extracellular region part, transcription factor activity, ion transport, and proteolysis. These pathways share certain genes, such as ADH4 and ALDH3B2. Previous studies have reported that xenobiotic metabolism by cytochrome P450 and drug activation are involved in tumorigenesis in RCC (38,39).

Collectively, the use of bioinformatic methods can significantly improve our understanding of RNA communication in ccRCC at the transcriptome level. However, these functional modules obtained from bioinformatic analyses require experimental confirmation.

In conclusion, our study represents the first comprehensive analysis of an lncRNA-miRNA-mRNA network in ccRCC. The results revealed certain novel lncRNAs involved in ccRCC, such as DGCR5, MYCNOS, and PART1, which may be involved in the progression of ccRCC through cytochrome P450-mediated drug metabolism. These findings expand the existing knowledge on lncRNA characteristics and provide new tools for the identification of lncRNAs in ccRCC, which can help to elucidate the disease processes and identify new targets for therapy.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

BL and TM performed the data analyses and wrote the manuscript. QL, SW, WS and WL contributed significantly in data analysis and manuscript revision. JL and YG conceived and designed the study. All authors read and approved the final manuscript and agree to be accountable for all aspects of the research in ensuring that the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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