Long Noncoding RNA DIO3OS Hinders Cell Malignant Behaviors of Hepatocellular Carcinoma Cells Through the microRNA-328/Hhip Axis

Zhanpeng Wang, Lina Song, Yanshuo Ye, Wei Li

Department of Hepatobiliary-Pancreatic Surgery, China-Japan Union Hospital of Jilin University, Changchun 130000, People’s Republic of China

Background: The decline of a long non-coding RNA (lncRNA) DIO3OS was implicated in a plethora of cancers, while the relevance in hepatocellular carcinoma (HCC) has not been mentioned. Accordingly, we set to determine the functional role of DIO3OS and the molecular mechanism in HCC progression.

Materials and Methods: The differentially expressed lncRNAs, mRNAs, and microRNAs (miRNAs) were obtained through the datasets GSE101728 and GSE57555. Afterwards, DIO3OS was enhanced in HCC cells to examine the behavior changes. Subcellular localization of DIO3OS was determined through website prediction and experimental validation. The expression of Hedgehog (Hh) signaling pathway-related genes was detected. The effects of DIO3OS overexpression on tumor growth were evaluated as well.

Results: DIO3OS was lower in HCC tissues and cells, while upregulation of DIO3OS repressed malignant biological behavior both in vitro and in vivo. DIO3OS, localized in the cytoplasm, inhibited the occurrence of HCC by disrupting the Hh pathway by sponging miR-328 to mediate Hh interacting protein (Hhip).

Conclusion: All in all, the obtained data suggested that DIO3OS interacted with Hhip-dependent Hh signaling pathway to inhibit HCC progression through binding to miR-328, which may be a potent therapeutic target for HCC.

Keywords: hepatocellular carcinoma, DIO3OS, competing endogenous RNA, microRNA-328, Hhip, hedgehog signaling pathway

Introduction

Liver cancer ranked the sixth most common carcinoma and the fourth most frequent consequence of cancer-associated death in 2018, among which hepatocellular carcinoma (HCC) contributed to 75%-85% of all cases diagnosed. The most important risk factor for the majority of HCC is the chronic liver disease caused from viral hepatitis, alcohol over-consumption, and/or nonalcoholic steatohepatitis. If diagnosed at an early stage, patients may be curable via resection, liver transplantation, or ablation to achieve an over 50% 5-year survival rate. Since HCC is recurrently resistant to chemotherapy and radiotherapy, many adjuvant therapies for HCC, such as small molecular targets, monoclonal antibodies and microRNA (miRNAs) have emerged. Long non-coding RNAs (lncRNAs) have been implicated in the pathogenesis and progression of various cancers, including HCC, and certain lncRNAs may function as diagnostic and/or prognostic markers for HCC.
Type 3 deiodinase (DIO3) and DIO3 opposite strand (DIO3OS) genes located at the distal end of the GTL2/DIO3 imprinted cluster are preferentially expressed in the GTL2/DIO3 imprinted region of mouse and humans. DIO3OS was remarkably reduced in patients with ulcerative colitis relative to normal controls. While its role in cancers, especially HCC remains elusive. Interestingly, IncRNAs may regulate the expression of microRNAs (miRNAs) which are closely related to the progression of HCC and have the potency to bind to IncRNA sponges to suppress gene expression and protein synthesis. miR-328 expression was promoted in 48 HCC tissue samples, and downregulation of miR-328 in HepG2 cells attenuated cell invasion and proliferation and promoted cell apoptosis. Moreover, circRNA-5692 bound to miR-328-5p which potentiated the malignant behaviors of HCC cells. Thus, we proposed that DIO3OS may interact with miR-328 to revert the malignant aggressiveness of HCC cells. More recently, hedgehog-interacting protein antisense RNA 1, a novel IncRNA, was monitored to interact with and stabilize the mRNA of hedgehog-interacting protein (Hhip) to enhance Hhip expression, thereby inhibiting the progression of HCC. In addition to facilitating development and regeneration, the Hedgehog (Hh) pathway results in alteration of progenitors when incessantly evoked, which might lead to small-cell lung carcinomas, prostate and digestive tract tumors. Considering the aforementioned investigations, in this work, we elaborated the probable roles of DIO3OS and miR-328 in HCC progression with the intention of offering insights for the treatment of HCC.

Materials and Methods

Subjects

We collected 31 tumor tissues and matched paracancerous tissues from patients with HCC diagnosed in China-Japan Union Hospital of Jilin University from April 2018 to July 2019, of which 19 were male and 12 were female, with a median age of 59.7 years and an average age of 56.31 ± 8.3 years. All patients were followed up every three months for a period of five years. All patients with clinical samples were diagnosed through pathological examination and none of them received radiotherapy or chemotherapy before enrollment. Patients complicated with chronic systemic diseases or other malignancies were excluded. All the samples were collected with written informed consent from patients, and the protocol was permitted by the ethics committee of China-Japan Union Hospital of Jilin University.

Bioinformatics Analysis

HCC-related datasets involving mRNAs and lncRNAs (GSE101728) and miRNAs (GSE57555) were found in the GEO database (https://www.ncbi.nlm.nih.gov/geo/). Differential expression of microarray expression matrix was conducted using in R-language limma package (http://master.bioconductor.org/packages/release/bioc/html/limma.html). The differentially expressed genes were screened out with the conditions of adj.p.Val < 0.05 and | LogFoldChange| > 1.5, and the R language pheatmap package (https://cran.r-project.org/web/packages/pheatmap/index.html) was applied for the heatmap plotting. The expression of DIO3OS and Hhip in The Cancer Genome Atlas (TCGA)-liver HCC database were then predicted with the help of GEPIA website (http://gepia.cancer-pku.cn/index.html). The binding miRNAs of DIO3OS and Hhip were predicted by StarBase, and screened by the jvenn website (http://jvenn.toulouse.inra.fr/app/example.html) for mapping.

Construction of Overexpression or Silencing Plasmids

The overexpression plasmid of DIO3OS, the corresponding empty vector (EV), two small interfering RNAs (siRNAs) targeting DIO3OS and corresponding scramble siRNA were synthesized by GenePharm Bioengineering Co., Ltd. (Shanghai, China).

Cell Treatment and Grouping

Human immortalized normal stem cell LO2 and HCC cell lines MHCC97, BEL-7402, HepG2, SMMC7721, and BEL-7405 cells were from the Cell Bank of Shanghai Institute of Cells, Chinese Academy of Science (Shanghai, China). All cells were cultivated in Roswell Park Memorial Institute-1640 medium supplemented with 10% fetal bovine serum (Gibco, Carlsbad, CA, USA). Subsequently, the well-grown BEL-7405 cells and HepG2 cells were transfected with constructed DIO3OS overexpression plasmid and the corresponding empty plasmid, respectively. The LO2 cells were transfected with siRNAs targeting DIO3OS and corresponding scramble siRNA. Transfection was performed using the lipofectamine 2000 kit (Thermo Fisher Scientific Inc., Waltham, MA, USA) strictly in accordance with the instructions for 48 h. Cells were harvested for following experiments. RT-qPCR was applied to examine DIO3OS expression to evaluate the transfection efficiency.
RT-qPCR
The total RNA in the tissues and cells was obtained using a TRizol kit (Thermo Fisher Scientific). The TaqMan miRNA reverse transcription kit (Applied Biosystems, Foster City, CA, USA) was used for reverse transcription of miRNA, while reverse transcription reagents (Applied Biosystems) were applied for mRNA and lncRNA reverse transcription. SYBR Green (Takara Bio Inc., Japan) was applied for real-time quantitative PCR. To synthesize cDNA, the reaction mixture was maintained at 16°C for 30 min, at 42°C for 30 min, at 85°C for 5 min, and then at 4°C. Next, 0.67 μL cDNA was amplified using 5 μL TaqMan 2× Universal PCR Master Mix (Applied Biosystems) without AmpErase Ung. 0.5 μL genespecific primers/probes, and 3.83 μL ddH2O. Quantitative PCR was performed on the StepOnePlus PCR system (Applied Biosystems), and the reaction mixture was incubated at 95°C for 10 min, followed by 40 cycles at 95°C for 15 s, followed by 1 min at 60°C. StepOne software v2.0 (Applied Biosystems) was applied to calculate the cycle threshold (Ct). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA was used as internal reference. The sequences were listed below: DIO3OS forward: AGAGTGGCACACTCATGTACCA, reverse: CCAAGT GCGCAACTCAGACAC; GAPDH: forward: CGTGAA AGATGACCCAGATCA; reverse: CACAGCCT GGATGGCTACGT.

Cell Viability Assays
The viability of HCC cells was detected by a cell counting kit-8 (CCK-8, Roche, Sweden) following the manufacturer’s instructions. Then, HCC cells were subjected to EdU staining with an EdU proliferation kit (ab219801, Abcam, Shanghai, China) for the proliferation activity measurement strictly according to a previous literature. Briefly, 1 × 10^6 cells were cultured with EdU for 4 h, followed by harvesting and fixation with 4% formaldehyde. Afterwards, EdU labeled cells were analyzed using a MoFlo Astrios (Beckman Coulter, Brea, CA, USA).

Detection of Apoptosis
Firstly, the propidium iodide (PI)/Annexin-V kit was used to label the cells in each group, and then the number of apoptotic cells was detected by a flow cytometer (Thermo Fisher). In short, the medium was discarded from each well, and cells were carefully washed with phosphate buffered saline (PBS) and preserved on ice. Cells were then detached with trypsin and isolated with 0.5 mL 0.5% trypsin. After that, they were resuspended in the original medium to insert isolated cells into the suspension for further examination. After exposure of 1.0 × 10^6 cells to 10 mL binding buffer and 1.25 mL Annexin V-fluorescein isothiocyanate, the cells were incubated at room temperature for 15 min in the dark. Then, the cell suspension was centrifuged at 1000 ×g for 5 min to remove the supernatant, and the cell was resuspended in 0.5 mL ice-cold 1× binding buffer. Finally, the cells were incubated with 10 mL propidium iodide and transferred to a fluorescent activated cell sorter tube (Fahrenheit, Munich, Germany). Fluorescence data analysis was performed via WinMDI 2.8 software.

Apoptotic cell levels were detected using a Hoechst 33258 staining kit (Roche) in strict accordance with the instructions. In short, Hoechst 33258 could bind to DNA molecules as fluorescent probes. The intake of Hoechst 33258 in apoptotic cells increased, and apoptotic cells showed strong blue fluorescence. After cell transfection, cells were fixed at 4°C for 10 min with 4% paraformaldehyde in 0.1 M PBS (pH = 7.4). Apoptotic cells were examined using a fluorescence microscope after staining with 5 mg/mL Hoechst 33258 for 10 min.

Transwell Assay for Cell Migration and Invasion
Transwell assay was fulfilled following a previous report. Serum-free medium was used to dilute Matrigel (BD Biosciences, San Jose, CA, USA) at a ratio of 6:1, and 100 μL diluted Matrigel was placed in the apical chamber at 37°C for 4 h for solidification. The apical chamber was filled with 100 μL cell suspension (1 × 10^5 cells), while the basolateral chamber was filled with 500 μL serum-free medium. After a night, noninvasive cells were removed with cotton swabs, and invaded cells on the surface of basolateral chamber were fixed in 4% paraformaldehyde for 30 min and stained with 0.1% crystal violet for 20 min. Finally, the invasive cells were visualized and calculated under a microscope. In terms of migration capacity, the steps were consistent except for Matrigel processing.

Sublocalization of DIO3OS
First, we used Lncatlas (http://lncatlas.crg.eu/) to predict the cellular sublocalization of DIO3OS. The localization of DIO3OS in HepG2 and BEL-7405 cells was then detected using a nuclear separation kit (PARIS kit; cat. no. AM1921; Invitrogen Inc., Carlsbad, CA, USA) in accordance with the
instructions. Subsequently, the positioning of DIO3OS in HepG2 and BEL7405 cells was further determined by fluorescent in situ hybridization (FISH) experiments, all of which were carried out as previously illustrated. In short, nucleic acid-modified oligonucleotide probes targeting DIO3OS (Exiqon, Vedbaek, Denmark) was used. 4′,6-diamidino-2-phenylindole (DAPI) was used to label nuclear DNA (in blue). The RNA signal was detected by incubation with biotinylated coupled antibody against digoxigenin, and amplified using SABC-Alexa Fluro 448 (in red).

Luciferase Reporter Assay
The binding relationships of miR-328 to DIO3OS and Hhip were detected using a luciferase reporter assay. DIO3OS or Hhip wild-type (WT) and 3’untranslated region (3’UTR) binding sequence mutant (MT) was synthesized by GenePharm Bioengineering Co., Ltd. (Shanghai, China) and inserted into pMIR-REPORT™ (Thermo Fisher Scientific). Luciferase activity was evaluated by a dual-luciferase reporter assay system (Promega Corporation).

Immunoblotting
Immunoblotting was exercised to examine the expression of Hh signaling pathway-related markers in HCC cells according to standard protocols as described previously. Protein was harvested using radio immunoprecipitation assay lysis buffer, and a bicinchoninic acid assay was applied to determine the protein concentration. Next, we separated equal amount of protein using 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred the protein to a polyvinylidene fluoride membrane. After blocking with 1.5% skimmed milk for 1 h at room temperature, membranes were incubated with the primary antibodies at 4°C overnight and with horseradish peroxidase-conjugated secondary antibodies. Membranes were developed using a chemiluminescent substrate. The antibodies were used as follows: Gli1 (1:200, ab49314, Abcam), Gli2 antibodies (1:500, ab26056) and Gli3 antibodies (1:500, ab6050) and the secondary antibodies (1:50,000, ab7090).

Tumorigenesis Assay in vivo
We subcutaneously injected $2 \times 10^6$ stably overexpressed HepG2 and BEL-7405 cells (suspended in 200 μL PBS) into the right axils of nude mice. We recorded tumor growth weekly and used the formula (tumor length $\times$ tumor width $\times$ tumor width $\times$ 1/2) to calculate tumor volume. After euthanasia (injection of pentobarbital sodium at 100 mg/kg), we weighed the tumor. We handled the experimental animals according to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health. The protocol was reviewed and approved by the Ethics Committee of China-Japan Union Hospital of Jilin University.

Immunohistochemistry
Tumor tissues were fixed with formalin, embedded in paraffin and cut into 5 μm thickness sections. The samples underwent strict processes of deparaffinization, rehydration, antigen recovery and endogenous peroxidase inhibition. Next, sections were immunoblotted overnight with the KI67 antibody (ab15580, Abcam) at 4°C and with the appropriate secondary antibody. As a chromogen, diaminobenzidine solution was applied for visualization. At last, nuclei were stained with hematoxylin. A microscope (Leica DM 4000B, Leica, Bannockburn, IL, USA) was used to acquire images.

Quantification and Statistical Analysis
SPSS 21.0 software (SPSS, Chicago, IL, USA) was used for statistical analyses. Normal distribution was tested by Kolmogorov-Smirnov. Results were displayed as mean ± standard deviation (SD). One-way or two-way analysis of variance (ANOVA) following Tukey’s post hoc test were performed to exhibit the differences among experimental groups. A level of $p < 0.05$ was considered significant.

Results
Poor Expression of DIO3OS Is Identified in HCC Patients and HCC Cell Lines
In the GEO database, we analyzed the GSE101728 dataset which included the cancer tissues of seven HCC patients as well as the paired adjacent tissues. Differential analysis of GSE101728 dataset was performed to screen out 30 significantly differentially expressed lncRNAs and to plot the heatmap, as shown in Figure 1A. Additionally, the expression of DIO3OS in HCC patients was analyzed in the TCGA database through the GEPIA bioinformatics website, which revealed the downregulation of DIO3OS in HCC patients (Figure 1B). Later, we tested DIO3OS in tumor tissues and paracancerous tissues of 31 HCC patients by RT-qPCR. The expression of DIO3OS in HCC tissues was found to be significantly reduced (Figure 1C). DIO3OS expression in HCC cells and LO2 cells was examined afterwards. As expected, DIO3OS was diminished in HCC cells (Figure 1D). With the aim to further verify the effect of DIO3OS on HCC, we transfected the DIO3OS overexpression plasmid into HepG2 as well as BEL-7405.
cells, whereas two siRNAs targeting DIO3OS into LO2 cells. RT-qPCR was then used to verify the transfection efficiency, and the expression of DIO3OS was significantly enhanced after overexpression plasmid delivery, while downregulated in LO2 cells following introduction of siRNAs (Figure 1E).

**DIO3OS Inhibits Malignant Behaviors in HepG2 and BEL-7405 Cells**

We found that after overexpression of DIO3OS, the number of EdU-positive cells was significantly reduced (Figure 2A), and the activity of HepG2 and BEL-7405 cells was significantly inhibited (as revealed by CCK-8) (Figure 2B). We further observed that restoration of DIO3OS resulted in promotions in HepG2 and BEL-7405 cell apoptosis (Figure 2C and D). On top of that, transwell assay unveiled the identical tendency as findings from CCK-8 and EdU assays. Resumption of DIO3OS hampered HepG2 and BEL-7405 cell invasion and migration (Figure 2E and F).

**DIO3OS Knockdown Promotes the Malignant Behaviors in LO2 Cells**

EdU staining and CCK-8 assays were then used to detect cell activity, we found that the proliferation of LO2 cells increased significantly after DIO3OS knockdown.
Moreover, the results of flow cytometry and Hoechst 33258 staining demonstrated that the apoptotic LO2 cells decreased remarkably after DIO3OS knockdown (Figure 3C and D). Finally, we showed through Transwell assays that DIO3OS knockdown contributed to increased number of migrated and invaded cells (Figure 3E).

**Figure 2** DIO3OS inhibits HepG2 and BEL-7405 cell malignant behavior. (A) EdU staining of proliferating cells. (B) HepG2 and BEL-7405 cell viability examined by CCK-8 assay. (C) apoptosis index of HepG2 and BEL-7405 cells examined by Hoechst 33258 staining. (D) PI/Annexin-V stained HepG2 and BEL-7405 cells determined by flow cytometry. (E) migration ability determined by transwell assays. (F) invasion ability determined by transwell assays. One-way ANOVA and Tukey’s multiple comparison test was applied to assess statistical significance. *p < 0.05.

(Figure 3A and B). Moreover, the results of flow cytometry and Hoechst 33258 staining demonstrated that the apoptotic LO2 cells decreased remarkably after DIO3OS knockdown (Figure 3C and D). Finally, we showed through Transwell assays that DIO3OS knockdown contributed to increased number of migrated and invaded cells (Figure 3E).

**DIO3OS Is Localized in Cytoplasm**

To this end, we predicted the main localization of DIO3OS with cytostasis (Figure 4A) through the Lncatlas bioinformatics prediction website. Furthermore, FISH experiments and nuclear separation experiments exhibited that DIO3OS mainly expressed in the cytoplasm in HepG2 and BEL-7405 cells (Figure 4B and C).
DIO3OS Serves as a ceRNA for miR-328 and Elevates Hhip Expression

Since DIO3OS was localized in the cytoplasm, it can be preliminarily demonstrated that DIO3OS played a role in stimulating the malignant aggressiveness of HCC cells through the ceRNA mechanism. Consequently, by analyzing the HCC-related miRNA dataset GSE57555, we found that there were 14 miRNAs with significant high expression in HCC. In addition, we used RNA22 to predict the miRNAs that might bound to DIO3OS, and compared the miRNA prediction results with the up-regulated miRNAs in the GSE57555 dataset. Only one miRNA was observed in the intersection: hsa-miR-328 (Figure 5A). It has been reported that downregulation of miR-328 inhibited metastasis and growth of HCC. Therefore, we used a luciferase activity assay to authenticate the relationship between miR-328 and DIO3OS (Figure 5B). We next examined miR-328 expression in HepG2 and BEL-7405 cells in the presence of DIO3OS, and found that DIO3OS diminished miR-328 expression (Figure 5C). Furthermore, we examined the miR-328 expression in tumor tissues and paracancerous tissues of 31 HCC patients, and found that miR-328 was higher in tumor tissues. The relationship between the expression of miR-328 and DIO3OS was analyzed by Pearson’s correlation. miR-328 was revealed to be negatively correlated with DIO3OS (Figure 5D and E).

Subsequently, we predicted the direct targets of miR-328 in RNA22, mirDIP, miRDB, DIANA and Targetscan, and compared the predicted results of target genes with those with poor expression in GSE101728 dataset. There were 2 intersection genes Hhip and DLG5 (Figure 5F). We analyzed the expression of Hhip in HCC patients in TCGA database through the GEPIA bioinformatics website. Hhip was significantly reduced in HCC patients (Figure 5G). In addition, Hhip was hypermethylated and transcriptionally reduced in HCC tissues and cells. Therefore, we used dual luciferase reporter gene assay to verify the binding relationship between miR-328 and Hhip (Figure 5H). We then examined the expression of Hhip in HepG2 and BEL-7405 cells overexpressing DIO3OS and found that overexpression of DIO3OS simultaneously increased Hhip expression (Figure 5I). Furthermore, we tested Hhip expression in 31 patients with HCC and in paracancerous tissues, and noted that Hhip expression was significantly lower in tumor tissues (Figure 5J). We lastly analyzed the relationship between Hhip and miR-328 or DIO3OS expression by Pearson’s correlation. Hhip was positively correlated with DIO3OS and negatively correlated with miR-328 (Figure 5K and L).

Figure 3 Silencing DIO3OS promotes LO2 cell malignant behavior. (A) EdU staining of proliferating LO2 cells. (B) LO2 cell viability examined by CCK-8 assay. (C) apoptosis index of LO2 cells examined by Hoechst 33258 staining. (D) PI/Annexin-V stained LO2 cells determined by flow cytometry. (E) migration and invasion ability determined by transwell assays. Unpaired t test was applied to assess statistical significance. *p < 0.05; **p < 0.01.
DIO3OS Disrupts the Hhip Signaling Pathway

It has been reported that Hhip inhibited proliferation of adipocytes through the Hh signaling pathway. Accordingly, the expression patterns of Hh signaling pathway-related proteins Gli1, Gli2 and Gli3 in HepG2 and BEL-7405 cells were evaluated. The expression of Gli1, Gli2 and Gli3 was decreased significantly following overexpression of DIO3OS (Figure 6A and B).

DIO3OS Inhibits HCC Growth in vivo

HepG2 or BEL-7405 cells with stable overexpression of DIO3OS were injected into the armpit of nude mice to evaluate the effect of DIO3OS on the growth of HCC cells in vivo. We monitored that overexpression of DIO3OS inhibited the growth rate of HepG2 or BEL-7405 cells in vivo, and overexpression of DIO3OS significantly reduced tumor weight at the end of the experiment (Figure 7A and B). Subsequently, we performed KI67 immunohistochemical staining on the tumors and found that overexpression of DIO3OS lowered the rate of KI67 positive cells (Figure 7C) in the tumor tissues.

Discussion

China is one of the regions that own the greatest HCC incidence, while the lack of appropriate preemptive or therapeutic treatment regimens reinvigorated wide research against HCC so as to develop novel therapeutic strategies. Dysregulation
**Figure 5** DIO3OS serves as a ceRNA by competitively combining to miR-328 with Hhip. (A) Targeting miRNAs of DIO3OS in GSE57555 dataset containing 6 HCC tissues and paired adjacent tissues predicted by RNA22. (B) The DIO3OS and miR-328 combination examined by dual luciferase assays. (C) miR-328 expression determined in HepG2 and BEL-7405 cells. (D) miR-328 expression in tumor and paracancerous tissues determined by RT-qPCR. (E) The association between miR-328 and DIO3OS analyzed by Pearson’s correlation analysis. (F) Targeting mRNAs of miR-328 in GSE101728 dataset. (G) Hhip expression examined by GEPIA website. (H) Hhip and miR-328 combination evaluated by dual luciferase assays. (I) miR-328 expression determined in HepG2 and BEL-7405 cells. (J) Hhip mRNA expression in tumor and paracancerous tissues determined by RT-qPCR. (K) the association between DIO3OS and Hhip analyzed by Pearson’s correlation analysis. (L) the association between miR-328 and Hhip analyzed by Pearson’s correlation analysis. The data are expressed as the mean ± SD. One-way ANOVA and Tukey’s multiple comparison test was used to determine statistical significance, *p < 0.05, **p < 0.01.

**Figure 6** DIO3OS inhibits the Hh pathway. (A) The protein expression of Gli1, Gli2 and Gli3 normalized to β-actin evaluated by Western blot assays. (B) The statistical analysis of panel A. The data are expressed as the mean ± SD. Two-way ANOVA and Tukey’s multiple comparison test was used to determine statistical significance, *p < 0.05.
Nevertheless, the relevance of We monitored that introduc-
Additionally, over-
In HCC, resumption of
Hence, to sightsee the
Of note, Gli3, a target gene
Then, we believed that the Hh pathway lies
invasive breast cancer,
non-small cell lung
cancer, invasive breast cancer,
non-small cell lung
cancer, and most importantly, HCC.
Additionally, over-
expression of metastasis associated in lung adenocarcinoma
tumor growth in vivo via performing as a ceRNA for miR-328
and upregulating Hhip expression through the Hh signaling
pathway. Altogether, these observations provided important
insights into the cancer-suppressing effect of DIO3OS in
HCC cell malignant aggressiveness.
In HCC, we discovered a relatively low expression of
DIO3OS in HCC tissue samples by accessing to the TCGA
and GEO databases, and RT-qPCR analysis exposed a similar
pattern in collected HCC tissues and cell lines. Moreover,
upregulation of DIO3OS through delivery of overexpression
plasmid of DIO3OS into HepG2 and BEL-7405 resulted in
decreases in growth, migration and invasion of these two
cell lines, while enhancements in cell apoptosis. By contrast,
DIO3OS knockdown by siRNAs led to the opposite trends in
LO2 cells. Recently, non-coding RNAs, including miRNAs
and lncRNAs, has been observed to produce a complex mod-
ulatory network through competing with each other to interact
with mRNAs, working as a ceRNA. Hence, to sightsee the
mechanism in which DIO3OS acts in HCC, we investigated
candidate miRNAs and established that DIO3OS competed
with miR-328 to slow down the progression of HCC. By
contrast, DIO3OS promoted cell proliferation and invasion
through competing for miR-122 to upregulate ALDOA
expression in pancreatic cancer. A plethora of researches on
miR-328 have paid much attention to its functions on cell
growth, invasion and migration in cancers, including gastric
cancer, invasive breast cancer, non-small cell lung
cancer, and most importantly, HCC.
Furthermore, in view of the mechanism of lncRNA-
mRNA-mRNA network, we screened out one target gene
of miR-328, Hhip. The level of Hhip was reduced in adipose
tissue development, and recombinant Hhip promoted
the differentiation of adipocytes by targeting the canonical
Hh pathway. More specifically, Hhip, a reverse modulator of the Hh pathway, was downregu-
lated in gastric cancer, and its overexpression hampered
cancer cell survival, proliferation, migration as well as
invasion. Then, we believed that the Hh pathway lies
the downstream of the DIO3OS/miR-328/Hhip axis in
HCC cells. The regulatory role of Hh pathway in liver
cells including hepatic stellate cells, progenitors, and
endothelial cells has been increasingly uncovered. Hh
pathway components (Gli1 and Gli2) were potentiated in
both HCC tissues and cells. We monitored that introduc-
tion of DIO3OS remarkably diminished the Hh pathway-
related proteins Gli1, Gli2 and Gli3 expression, while
restored Hhip expression. The observations of our study
were largely in agreement with a previous study, where the
lncRNA-dependent noncanonical Hh pathway in breast
cancer sheds some lights on certain patients who would
respond optimally to treatment. In HCC, resumption of
the Hh pathway is accountable for the maintenance of
cancer stem cells, and Hh disruption demonstrates an
obvious anti-tumor effect. Of note, Gli3, a target gene

Figure 7 DIO3OS inhibits HCC tumor growth in vivo. HepG2 and BEL-7405 cells stably overexpressing DIO3OS were inoculated subcutaneously into BALB/c nude mice at a dose of 2 × 10⁶ per mouse (n = 6 in each group). Tumor growth was measured continuously every 7 days. At 35 days post-implantation, the mice were euthanized by pentobarbital sodium at 100 mg/kg. (A) growth curve for the tumor size of nude mice. (B) the tumor weights of nude mice. Tumor sections were obtained and stained with antibody against KI67. (C) representative images of KI67-positive tumor cells and quantification of immunostaining. In panel B and C, one-way ANOVA, while in panel A, two-way ANOVA and Tukey’s multiple comparison test was used to determine statistical significance, * p < 0.05, ** p < 0.01.
for miR-133, was repressed by miR-133 during the event of embryonic myogenesis.\textsuperscript{35} However, the correlation between miR-328 and the Hh pathway in HCC cells was not investigated in this work due to the time and fund limitations.

## Conclusion
Lastly to generalize our finding, we recapitulate a DIO3OS/miR-328/Hhip regulatory network that might regulate the HCC cell growth, migration and invasion. Meanwhile, DIO3OS and miR-328 procedure a regulatory loop where the release of DIO3OS contributes to the declines in miR-328 expression, eventually enhancing Hhip expression and disrupting the Hh pathway. Thus, this study may provide a new therapeutic maneuver for HCC treatment and offer some insights on the lncRNA-miRNA-mRNA regulatory nexus on the HCC pathogenesis.

## Author Contributions
ZPW contributed to the study design and manuscript preparation; All authors contributed to data analysis, drafting or revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

## Disclosure
The authors declare no conflict of interest.

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