Deciphering The Molecular Mechanism of Long Non-Coding RNA HIFA-AS1 Regulating Pancreatic Cancer Cells

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Research Article

Keywords: pancreatic cancer, long non coding RNA, HIFA AS1 , overexpression, molecular mechanism

DOI: https://doi.org/10.21203/rs.3.rs-533849/v1

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Deciphering the molecular mechanism of long non-coding RNA HIFA-AS1 regulating pancreatic cancer cells

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Abstract

Background: *HIF A-AS1*, an antisense transcript of *HIF1α* gene, is a 652-bp long non-coding RNA (lncRNA) which globally expressed in multiple tissues of animals. Recent evidence indicated that the *HIF A-AS1* was involved in tumorigenesis of several types of cancer, but there were no reports on pancreatic cancer (PC).

Results: In order to investigate whether the *HIF A-AS1* could mediate the PC or not, it was overexpressed in a PC cell line (PANC-1), and a series of experiments including cell viability detection, flow cytometry, transwell migration, clone formation and wound healing were performed. Functionally, the results indicated that overexpression (OE) of *HIF A-AS1* could inhibit proliferation and shift, and promote apoptosis of PC cells. Moreover, to explore underlying molecular mechanism of anti-tumorigenic actions of *HIF A-AS1* in PC cells, the iTRAQ (isobaric tags for relative and absolute quantification) quantitative proteomics analysis was implemented and the results indicated that OE of *HIF A-AS1* globally affected the expression levels of multiple protein associated with metabolism of cancer. Moreover, the network analysis revealed that the most of these differentially expressed proteins (DEPs) were integrated, and severed essential roles in regulatory function.

Conclusions: In summary, *HIF A-AS1* may exhibit a potential therapeutic effect on PC, and our study provided useful information in this filed.

Keywords: pancreatic cancer, long non-coding RNA, *HIF A-AS1*, overexpression, molecular mechanism
Introduction

Pancreatic cancer (PC) remains one of the most common causes of cancer-related mortality [1] at the seventh in humans worldwide [2], with 5-year overall survival rate of less than 5% [3]. In most cases, PC develops with usually clinically silent at the early stage, but the variable symptoms, local invasiveness, or metastases only develop at an advanced stage [4]. Nowadays, the therapeutic efficacy of PC treatment is still very limited, and far from satisfactory [5, 6]. Hence, in order to enhance the cure rate of PC, it is necessary to investigate the molecular mechanisms, which would provide new opportunities to improve effective therapeutic strategies against PC.

Long non-coding RNAs (lncRNAs), a kind of non-coding RNAs transcripts, comprises longer than 200 bp without protein-coding potential [7-9]. Current studies have showed that lncRNAs could mediate gene expression via chromosome remodeling, transcription and post-transcriptional processes [10]. As so far, increasing evidences demonstrate that lncRNAs play an important role in regulating vital molecular mechanism [11] and biological functions of the cells [12, 13], such as proliferation, migration, invasion, cell cycle and apoptosis [14-16]. Without a doubt, various expression of lncRNAs could contribute to tumor development and progression [17], but its regulatory mechanism had not been completely investigated.

HIF1A-AS1 is an antisense transcript of HIF1α [18], and accumulating evidence has revealed that it plays a key role in proliferation and apoptosis of vascular smooth muscle cells [19-21], and human hepatic stellate cells [22]. Furthermore, it promotes tumor necrosis factor-α-induced apoptosis [23], thereby affecting the occurrence and
development of thoracic aortic aneurysm [24]. HIF1A-AS1 can regulate starvation-induced hepatocellular carcinoma cell apoptosis, promoting hepatocellular carcinoma (HCC) cell progression [25]. Therefore, HIFA-ASI has the capacities to affect the occurrence and development of multiple types of cancer, but there is no report on the molecular regulation mechanism of HIFA-ASI in PC.

In the current study, to explore whether HIFA-ASI could regulate PC or not, we constructed overexpression (OE) plasmids containing HIFA-ASI, and transferred them to a PC cell line. Subsequently, a series of experiments, including cell viability detection, flow cytometry, transwell migration, clone formation and would healing were conducted. And the experimental results showed that OE of HIFA-ASI could inhibit proliferation and metastasis, and promote apoptosis of PC cells, comparing with the normal control (NC) cells. In order to further explore the molecular mechanism of HIFA-ASI regulating PC cells, we collected cell samples from OE and NC groups for iTRAQ (isobaric tags for relative and absolute quantification) proteomics experiments. Here we report the results.

Materials and methods

Cell culture

The human pancreatic cancer cell line (PANC-1) was provided by Procell (Wuhan, China) and cultured in monolayers in Dulbecco's modified Eagle's medium (DMEM) (Invitrogen, Carlsbad, USA). All media were supplemented with 10% fetal bovine serum (Hyclone UT, USA) in the presence of 100 U/ml penicillin and 50 μg/m streptomycin (Beyotime, Shanghai, China) with humidified atmosphere of 5% CO₂ and 95% air at 37 °C.
Plasmid construction, Lentivirus package and transfection

*HIF1A-AS1* was cloned into the pcDNA3.1(+) vector using the restriction sites for *KpnI* (GGTACC) and *XhoI* (CTCGAG), and this 652 bp insert was verified by sequencing. Two μg of plasmids containing *HIF1A-AS1* were mixed with the lentivirus packaging plasmids pMDLg-pRRE, pMD2.G, and pRSV-Rev according to the previous standard protocol [26]. Subsequently, PANC-1 cells were infected with 20 multiplicity of infection (MOI) lentivirus for 24 h and incubated in fresh medium. The cells were washed with fresh complete media after 24 h and the efficiency OE of *HIF1A-AS1* was verified by quantitative RT-PCR (qRT-PCR).

RNA extraction and qRT-PCR

Total RNA was extracted from the cells using TRIzol reagent (Ambion, Austin, USA) and further purified with two phenol-chloroform treatments, then treated with RQ1 DNase (Promega, Madison, USA) to digest DNA. The quality and quantity of the purified RNAs were determined using a Nano Photometer spectrometer with the absorbance at 260 nm/280 nm and next were verified by 1.2% agarose gel electrophoresis. The cDNA was synthesized with random primers with the High-capacity cDNA Reverse-Transcription Kit (Takara, Dalian, China), and real time PCR was implemented for detecting gene expressions using the designed primers in Table 1 with SYBR Green I dye (Qiagen, Hilden, Germany). The PCR conditions were as follows: pre-denaturation at 95 °C for 1 min, 40 cycles of denaturing at 95 °C for 15 s, annealing at 60 °C for 30 s and elongation at 72 °C for 40 s. The relative expression of genes was analyzed by the $2^{-\Delta\Delta CT}$ method with the *Actin* as an internal control [27].

Cell viability detection

The viability of PANC-1 cell was evaluated using the CCK-8 assay (Solarbio, Peking,
China) according to the instructions of the manufacturer. The cells were slightly seeded into the 96-well plates with 100 μl suspension per well overnight. At 0, 24, 48 and 72 h, 10 μl of CCK-8 solution was added to each well, and then the plates were incubated for 0.5 h. At last, absorbance was measured at 450 nm by microplate reader (Bio-Rad, Hercules, USA).

Flow cytometric detection

The apoptosis of the PANC-1 cells were determined by flow cytometry with Annexin V-conjugated FITC Apoptosis detection kit (BD, Franklin Lakes, USA). After infection for 24 h, cells were harvested and washed twice with PBS, then re-suspended in 5μl FITC-conjugated anti-Annexin V antibody and in 500 μl binding buffer with 5μl Propidium iodide (PI). Apoptosis was measured with a FACS calibur flow cytometer MoFLO XDP (Beckman, Hercules, USA).

Transwell invasion

For the transwell assay, the properties of migration of cells were evaluated by using 24-well transwell plates (Corning, NY, USA). About 1×10^4 cells per well were seeded into the upper chamber with serum free medium in triplicate. Medium containing 10% FBS (300 μl) was added to the DMEM of 5% CO2 and 95% air at 37°C. After incubation for 24 h, the medium were removed, and cells were fixed with 4% paraformaldehyde for 15 min, stained with 0.1% crystal violet for 20 min, and counted from five randomly chosen fields for each well by stereo microscope (Leica, Wetzlar, Germany).

Wound Healing Assay

After 12 h of the transfected cells were seeded in 6-well plates, confluent monolayers in each well were washed with PBS and created using a 200 μl sterile pipette tip to
generate a wound. Wound healing was evaluated and photographed images were taken by 200 magnification a Zeiss microscope (Leica, Wetzlar, Germany) from each well at 0, 24, and 48 h post-injury time points after the wound was made.

**Cell clone formation assay**

PANC-1 cells were plated into 6-well plates (800 cells per well) and cultured for 10–14 days, then were digested at the logarithmic phase to make a single-cell suspension using culture medium. Cell were stained with 0.4% crystal violet (Bio Basic Inc., Markham, Canada). Finally, the number of colonies was calculated under an inverted microscope (Leica, Wetzlar, Germany).

**Western Blotting Analysis**

PANC-1 cells were collected and the homogenates were centrifuged for 30 min at 4 °C, 12,000 rpm with cell lysis buffer. Then the protein extracts were separated on 10% SDS-PAGE and transferred to polyvinylidene difluoride (PVDF) membranes. Membranes were incubated at room temperature for 1 h in a 5% skim milk TBST blocking solution and incubated with agitation at 4 °C overnight with specific primary antibodies anti-GAPDH (Beyotime, Shanghai, China), anti-Bax (Cell signaling technology, Beverly, MA, USA ), anti-P53 ( Szybio, Wuhan, China), anti-Caspase 3 (Cell Signaling Technology), and anti-PARP-1 (Cell Signaling Technology). Next, membranes also were incubated with secondary antibodies conjugated by horseradish peroxidase (HRP) (Zhong san jinqiao, Beijing, China) for 50 min at room temperature. At last, protein bands were determined using the Western blotting detection system (GE Healthcare, Amersham, UK).

**iTRAQ quantitative proteomics analysis**
For protein extraction from the cells (5×10^6), the lysis buffer (7M Urea/2M Thiourea/4% SDS/40 mM Tris-HCl, pH 8.5) that contains 1 mM phenylmethanesulfonyl fluoride (PMSF) and 2 mM ethylenediaminetetraacetic acid (final concentration) was added to the sample, mixed and incubated in ice for 5 min. Subsequently, DL-Dithiothreitol (DTT) was added to a final concentration of 10 mM. The lysate was sonicated on ice for 20 min, then centrifuged at 4 °C, 13,000 g for 25 min. The supernatant was mixed with four volumes of precooled acetone and kept at -20 °C overnight. After centrifugation, the protein pellets were air-dried and re-dissolved in 8M urea/100mM triethylamine borane (TEAB) (pH=8.0). Protein samples were reduced with 10 mM DTT at 56 °C for 35 min and alkylated with 50 mM iodoacetamide (IAM) in the dark at room temperature for 30 min. The supernatant was transferred to a new centrifuge tube, and the protein precipitation was performed by acetone precipitation. The protein pellet was re-dissolved by adding 8 M urea/100 mM TEAB (pH=8.0) solution, and DTT was added to a final concentration for 10 mM, with reduction reaction at 56 °C for 25 min. Subsequently, IAM was added to a final concentration for 55 mM, and the alkylation reaction was carried out at room temperature for 40 min in the dark. The protein concentration was assessed by the Bradford method.

The 100 μg protein were trypsin digested with trypsin each sample. After diluting the protein solution 5 times with 100 mM TEAB, trypsin were added by a mass ratio of 1:50 (trypsin: protein) overnight at 37 °C. The peptides were desalted with C_{18} column after enzymolysis, and the desalted peptides were vacuum freeze-dried.

The mass spectrometry data was collected and analyzed using the Eksigent nanoLC
system (SCIEX, USA) coupled to the TripleTOF 5600+ mass spectrometers. Samples were iTRAQ labeled as follows: NC-1, X1; NC-2, X2; NC-3, X3; OE-1, X4; OE2, X5; and OE-3, X6. And all of the labeled samples were mixed with equal amounts. Next, the polypeptide solution was added to analytical ChromXP C\textsubscript{18} column (Bonna-Agela Technologies Inc., Wilmington, DE) (5 μm, 100 Å, 4.6×250 mm), and eluted at 300 nl/min on a C\textsubscript{18} analytical column (3 μm, 75 μm× 150 mm) over 90 min gradient. The two mobile phases were used both buffer A (2% acetonitrile / 0.1% formic acid / 98% H\textsubscript{2}O) and buffer B (98% acetonitrile / 0.1% formic acid / 2% H\textsubscript{2}O). For Information Dependent Acquisition (IDA), the first-order mass spectrum was scanned with an ion accumulation for 250 ms, at the same time the secondary mass spectrum of 30 product ion scans were collected with 50 ms. The MS1 spectrum was acquired in the range 350-1500 m/z, and the MS2 spectrum was acquired in the range 100-1500 m/z. Precursor ions were set from reselection for 15 s.

The original MS/MS file data were analyzed using ProteinPilot Software v4.5 (AB Sciex, Shanghai, China). For protein identification, the Paragon algorithm, which was integrated into ProteinPilot, was used against the UniProt/SwissProt database for database searching. The parameters were set as follows: the instrument was TripleTOF 5600+, iTRAQ quantification, and cysteine modified with IAM, and biological modifications were selected as ID focus, trypsin digestion, quantitate, bias correction, and background correction was used for protein quantification and normalization. For calculation of the false discovery rate (FDR), an automatic decoy database search strategy was used to estimate FDR using the proteomics system performance evaluation pipeline software (PSPEP, integrated into the ProteinPilot Software). Unique peptides were used for iTRAQ labeling quantification, and peptides with global FDR values from fit less than 1% were considered for further analysis. Within each iTRAQ run,
differentially expressed proteins (DEPs) were determined based on the ratios of differently labeled proteins and p values provided by ProteinPilot, the p values were generated by ProteinPilot using the peptides used to quantify the respective protein. For the determination of DEPs, fold changes (FC) were calculated as the average comparison pairs among biological replicates. Proteins with FC larger than 1.2 and a p < 0.05 were considered to be changes that are significantly different.

**Bioinformatics and annotations**

The biological and functional properties of all the identified proteins were analyzed by matching to NCBI (http://www.ncbi.nlm.nih.gov/) and Swiss-Prot/UniProt (http://www.uniprot.org/) databases, and were mapped with Gene Ontology (GO, http://www.geneontology.org/) and the Cluster of Orthologous Groups of proteins (COGs, http://www.ncbi.nlm.nih.gov/COG/) databases. The pathway enrichment analysis and metabolic pathways about the identified proteins were annotated by Kyoto Encyclopedia of Genes and Genomes (KEGG) mapping (http://www.genome.jp/kegg/). STRING v10.1 (http://string-db.org/) was applied to explore and analyze the protein-protein interaction (PPI) information of DEPs to evaluate the interactive associations. The PPI network was constructed and visualized using Cytoscape software (version 3.5.1; www.cytoscape.org).

**Statistical analyses**

All data analysis was performed using the SPSS16.0 statistical software as the mean ± standard deviation. Statistically significant differences comparison between two groups means were analyzed by Student’s t-test. P value <0.05 was considered to be statistically significant.
Data availability statement

The datasets generated and/or analysed during the current study are available in the ProteomXchange repository (Accession No: IPX0003153000). For the interview, the datasets could also be obtained from a web link:

https://www.iprox.org/page/PSV023.html/?url=1622727859237GpMj, with a code: E5xT.

Results

HIFA-ASI regulates the apoptosis and proliferation of PC cells

To investigate the role of HIFA-ASI in the PC cells, an OE vector containing the HIFA-ASI was transfected into PANC-1 cells. The qRT-PCR experiment was applied to measure the efficiency of OE, and the results showed that expression levels of HIFA-ASI in OE cells was about 10000-fold more as compared with the normal control (NC) cells (Fig. 1A), demonstrating a successful establishment of human PC cells with OE of HIFA-ASI.

To explore whether the HIF-AS1 regulate the proliferation of PC cells or not, the experiments including CCK-8, were conducted for PANC-1 cells from OE and NC groups. The results indicated that viability cells with HIF-AS1 OE was declined significantly, during varying time periods (0, 24, 48 and 72h) (P<0.001) (Fig. 1B and C).

Furthermore, flow cytometry was used to determine whether HIF-AS1 could affect apoptosis of PANC-1 cells or not. It (Fig. 1D and E) displayed that the OE of HIF1A-ASI significantly promoted apoptosis of PC cells, and the number of apoptotic cells
obviously increased about 50% compared with the NC group (Fig. 1D). The above results indicated that HIF1A-AS1 had the capacities to regulate proliferation and apoptosis of the PC cells. Further, western blot analysis reported that the expression levels of Cleaved caspase-3, Bax, P53, and PARP-1A protein were higher in OE group (Fig. 1F), thereby inducing apoptosis in pancreatic cancer.

**HIF-A-S1 regulates the migration of PC cells**

To further explore the role of HIF1A-AS1 in regulating metastasis of PANC-1 cells, the transwell migration assays were performed. It showed that the migration ability of cells from OE group was reduced about 50% (P<0.001) (Fig. 2A and B). In terms of cell clone, the clones formed in NC had a greater cell number about 45% compared with OE (Fig. 2C and D). Cell migration was detected using Wound-healing assay in PANC-1 cells. The results from invasion assay showed that OE of HIF-AS1 promoted cell invasion after transfection for 24 and 48 hours (Fig. 2E). And the difference of the wound width after 24 hours of transfection is the most significant compared with the comparison, which exceeds the control by about 20% (Fig. 2F), suggesting a functional role for HIF1A-AS1 in inhibiting metastasis of the PC cells.

**The summary of iTRAQ proteomics analysis**

To explore the molecular mechanism of HIF1A-AS1 mediating the proliferation, apoptosis and shift of PANC-1 cells, an iTRAQ was applied to uncover altered protein expressions and signaling pathways.

In total, the quality of the data obtained from the iTRAQ was analyzed using parameters
such as coefficient of variation about repeatability, distribution of unique peptide, peptide length, and distribution of coverage (Table 2). First of all, for the repeatability, there is a little difference concentration of CV data between NC and OE groups, and the cumulative percentages of CV were 7.81% and 7.29% respectively, indicating that the PANC-1 samples in each group are more reproducible (Fig. 3A).

In accordance with unique peptide determined as the peptide identified only for one protein, the presence of the corresponding protein can be uniquely determined. Then for the distribution of unique peptide number, the two-coordinate distribution map showed the number of unique peptides contained in all the proteins identified in this assay. For example, when the x-axis, left y-axis and right y-axis are 2, 646 and 26.25 respectively, it means that there are 646 proteins with 2 as the unique number of peptides, which account for 26.25% of the total number of proteins obtained (Fig. 3B).

Subsequently, the length of the identified peptides was analyzed. The average length of the polypeptide was 11.56 and within a reasonable range. Moreover, the length of the identified peptides was mainly concentrated between 7 and 15, and 9 was the maximum number (Fig. 3C). In addition, the protein identification coverage could reflect the overall accuracy of the identification results indirectly. The different colored pie represented the percentage of proteins with different identification coverage ranges. It showed that 37.21% proteins were with the peptide coverage less than 10%, and 39.51% proteins had more than or equal to 20% of the peptide coverage, with the average protein identification coverage being 19.53% (Fig. 3D).
A total of 4872 proteins were identified in all samples, and 4738, 2475 and 2539 ones were annotated successfully by GO, COG and KEGG, respectively (Fig. 3E). Particularly, the GO enrichment for the 4738 annotated proteins was carried out, including cellular localization (CC) (Data not shown), molecular functions (MF) (Data not shown) and biological processes (BP). The BP classification showed that most of these proteins were enriched in cellular process (13.04%), metabolic process (11.26%), biological regulation (8.55%), regulation of biological process (8.13%), cellular component organization or biogenesis (7.00%) and so on (Fig. 3F).

**Exploration of DEPs and functional analysis**

On basis of the relative quantitative results, 338 DEPs were found in OE VS NC according to FC and p value (FC ≥ 1.2 or ≤ 0.83, p ≤ 0.05), and the up-regulated and down-regulated ones were 183 (Table 3) and 155 (Table 4), respectively. The protein abundance distribution graph, and the volcano plot showed the proportion of DEPs in the total identified proteins (Fig. 4A and 4B). A hierarchical clustering analysis of DEPs was also performed (Fig. 4C).

In addition, KEGG enrichment for DEPs was implemented. It can be seen from the pie chart that the top 10 pathways were different among all up-regulated and down-regulated proteins. KEGG pathway enrichment was also variable across the up-regulated proteins group and down-regulated proteins group. The enriched pathways for up-regulated proteins, include “RNA transport” (ID: ko03013), “Metabolic pathways” (ID: ko01100), “Ribosome” (ID: ko03010), “Spliceosome” (ID: ko03040), “Microbial metabolism in diverse environments” (ID: ko01120), “Pathogenic
Escherichia coli infection” (ko05130), “Protein processing in endoplasmic reticulum” (ko04141), “Purine metabolism” (ko00230), “Glycolysis/ Gluconeogenesis” (ko00010), “Focal adhesion” (ko04510) and so on.

Furthermore, down-regulated proteins were primary enriched in some pathways, including “Metabolic pathways” (ID: ko01100), “Protein processing in endoplasmic reticulum” (ID: ko04141), “Pathways in cancer” (ID: ko05200), “Arrhythmogenic right ventricular cardiomyopathy” (ID: ko05412), “Microbial metabolism in diverse environments” (ID: ko01120) and “Regulation of actin cytoskeleton” (ko04810), “Tight junction” (ko04530), “Hypertrophic cardiomyopathy (HCM)” (ko05410), “Peroxisome” (ko04146), “Dilated cardiomyopathy” (ko05414) and so on (Fig. 4D).

According to these results indicate that the inhibitory effects of $HIF1A-AS1$ on the proliferation, apoptosis and migration of PANC-1 cells may be related to its capability to regulate protein interactions, catalytic activity and enzyme regulator activity. The top 10 pathway metabolic function types were different in all up-regulated and down-regulated differential proteins by KEGG. Four types were same containing metabolic pathway, regulation of actin cytoskeleton, microbial metabolism in diverse environment, protein processing in endoplasmic reticulum. Pathway analysis revealed that “Metabolic pathways” at the second of up-regulated genes and at the first in down-regulated genes in the enrichment results. Moreover, OE of $HIF1A-AS1$ may exhibit anticancer effects by regulating the pathways associated with metabolism of cancer. Therefore these results suggested that $HIF1A-AS1$ might affect RNA polymerase to
control the transcription of downstream tumor-associated genes as to antagonize the proliferation, apoptosis and migration of PC cells.

**Construction of DEPs Protein-protein interaction (PPI) network**

PPI network of common DEPs was constructed by the STRING online database and Cytoscape software to analyze the interactions of DEPs because the String database could identify interactions between known proteins and predictive proteins (Fig. 5). A total of 338 DEPs (155 down-regulated and 183 up-regulated) were filtered into the DEPs PPI network complex. The wonderful network suggested that these DEPs might work together to regulate apoptosis, proliferation and invasion of PC cells. These proteins are expected to become targets for the treatment of PC.

**Expression Validation by qRT-PCR**

To confirm the veracity and reliability of the proteomic assays, the expression levels of five candidate proteins were measured by qRT-qPCR, including MX1 (Interferon-induced GTP-binding protein Mx1), IFIH1 (Interferon-induced helicase C domain-containing protein 1), IFIT1 (Interferon-induced protein with tetratricopeptide repeats 1), ISG15 (Ubiquitin-like protein ISG15), P4HB (Protein disulfide-isomerase), SOD2 (Superoxide dismutase [Mn], mitochondrial) (Fig. 6). Some specific primers were designed for these candidate (Table 1). MX1, IFIH1, IFIT1 and ISG15 mRNAs showed more than 2-fold down-regulation as compared to NC (Fig 6A, B, C and E) and P4HB more than 0.5- fold down-regulation (Fig. 6D). In addition, SOD2 showed 2 fold up-regulation (Fig. 6F). In short, above data supported that the results of the proteome was
Discussion

PC remains one of the deadliest cancer types and worlds’ most aggressive malignancies [28]. Accumulating reports have reported that the potential of IncRNAs as diagnostic or prognostic biomarkers ubiquitously dysregulated and have crucial regulatory roles in tumor cells, including PC [29]. However, the regulatory mechanisms of multiple IncRNAs are elusive in many kinds of cancers such as thoracic aortic aneurysm and HCC. Herein, we first aimed to explore the molecular mechanisim of HIF1A-AS1 regulating PC.

In the present study, we investigated the biological function of HIF1A-AS1 on proliferation, apoptosis, and metastasis of PC cells. Consistently, it was found that HIF1A-AS1 was a suppressor of cell growth and progression in PC. Firstly, up-regulation of HIF1A-AS1 inhibited cell growth and promoted apoptosis in PANC-1 cancer cells. Moreover, HIF1A-AS1 inhibited cell migration. Actually, the function of HIF1A-AS1 in other tumors has been reported. For instance, higher expression of HIF1A-AS1, a novel diagnostic predictor, could be clinically functioned as a potential biomarker in colorectal carcinoma [30]. Besides, the levels of HIF1A-AS1 were significantly increased in tumor tissues or serum from non-small cell lung cancer patients [31]. Above reports researched with clinical samples, but may be the opposite with this study using PC cells. Therefore, this study indicated HIF1A-AS1 as an important role for a novel mechanism in the PC modulated progress, could develop as a potential therapeutic target or biomarker for PC prevention and control.
The quantitative proteomics analysis revealed that the expression levels of \( SOD2 \) was up-regulated. Previous studies have revealed that \( SOD2 \) has both tumor suppressive and promoting functions, which are primarily related to its role as a mitochondrial superoxide scavenger and \( \text{H}_2\text{O}_2 \) regulator [32]. \( SOD2 \) is role as both a tumor suppressor in early tumorigenesis and as a tumor promoter during metastatic progression [33]. Therefore, \( HIF1A-AS1 \) could regulate \( SOD2 \) to inhibit the metabolic developmental process in PC. In addition, \( Mx1 \) expression was inversely correlated with prostate cancer [34]. However, in this study the expression levels of \( MX1 \) was positive correlation with PC. \( MX1 \) [35], the members of IFN-stimulated genes [36], \( HIF1A-AS1 \) could regulate \( MX1 \) of type I interferon-mediated signaling pathway to restrain PC. According to previous reports, \( IFIT1 \) [37], OE of \( IFIT1 \) involved in a variety of biological processes, such as cell proliferation, migration and tumor growth [38, 39].

In this review, we hypothesize that \( HIF1A-AS1 \) could inhibit \( IFIT1 \) to regulate the cell proliferation, migration to restrain PC to some extents. \( IFIH1 \), may play an important role in enhancing natural killer cell function and may be involved in growth inhibition and apoptosis in several tumor cell lines [40, 41]. In a word, the involvement of \( IFIH1 \) in the growth inhibition and apoptosis of PANC-1 cancer cells was regulated by \( HIF1A-AS1 \). \( ISG15 \), induce natural killer cell proliferation, act as a chemotactic factor for neutrophils and act as an IFN-gamma-inducing cytokine lays a substantial role in the antiviral state induced by IFN [42, 43]. Further, we showed that inhibition of PC by \( HIF1A-AS1 \) and \( ISG15 \) is sufficiently. In line with these studies, the present findings demonstrated that \( HIF1A-AS1 \) might regulate some tumorigenesis of PANC-1 cancer
cells via targeting interferon-mediated signaling pathway, ubiquitin system and H₂O₂ regulator all closely related to metabolic pathways in cancer. In addition, tumor-suppressive role \textit{HIF1A-AS1} positively regulated the expression of \textit{SOD2}, and negatively regulated the five gene expression related to metabolic regulation to suppress PC growth and progression.

In summary, these findings demonstrated that \textit{HIF1A-AS1} could inhibit cell growth and progression of PANC-1 cells. In the future studies about diagnostic specificity and sensitivity of \textit{HIF1A-AS1}, may be used as a potential biomarker and guidance for early diagnosis of PC. Although clinical applications need to be further explored, these results further provided insight into the molecular mechanisms associated with the tumorigenesis and scientific experimental basis for the treatment of PC.
Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and materials

The datasets generated and/or analysed during the current study are available in the ProteomXchange repository (Accession No: IPX0003153000). For the interview, the datasets could also be obtained from a web link:

https://www.iprox.org/page/PSV023.html?url=1622727859237GpMj, with a code: E5xT.

Competing interests

The authors declare that they have no competing interests.

Funding

Supported by Natural Science Funding of Liaoning Province (#2019-ms-350) and SYDW [2018]09 Laboratory Animals Project Funding of Military.

Author Contribution

He Zhang, Rui Sun and Jing Tian designed and managed the project. He Zhang, Xinwei Wang and Yue Wu drafted the manuscript. Xuehua Li, Rui Sun and Jing Tian participated in sample collection and manuscript revision. All authors read and approved the final manuscript.

Acknowledgement
Supported by Natural Science Funding of Liaoning Province (#2019-ms-350) and SYDW [2018]09 Laboratory Animals Project Funding of Military.
Figure 1. Overexpression (OE) of *HIF1A-AS1* affects the proliferation, apoptosis of the pancreatic cancer (PC) cells (Pan-1). (A) Real-time PCR showed that the levels of *HIF1A-AS1* was significantly increased in the cells from OE group compared with normal control (NC). (B) & (C) OE of *HIF1A-AS1* reduces significantly the viability of PC cells at 24, 48, and 72 h. (D) OE of *HIF1A-AS1* promotes significantly apoptosis of PC cells assessment of cellular apoptosis using Annexin V - fluorescein isothiocyanate staining coupled with flow cytometry. (E) Total percentage of apoptotic PANC-1 cells in each group are summarized with data presented as the mean ± SD of three independent experiments. (F) Western blotting reveal Cleaved caspase-3, Bax, P53, and PARP-1A protein expression in PANC-1 cells. ***P<0.001
Figure 2. Overexpression (OE) of *HIF1A-AS1* regulates metastasis of the pancreatic cancer (PC) cells (Pan-1). (A) & (B) OE of *HIF1A-AS1* inhibits metastasis of PC cells by transwell migration assay. (C) & (D) OE of *HIF1A-AS1* inhibits metastasis of PC cells by clone formation assays. (E) & (F) OE of *HIF1A-AS1* inhibits metastasis of PC cells by wound-healing assay. (n = 3 cultures, paired Student’s t-test, ± SD) ** P<0.01, *** P<0.001
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### Tables and Table Legends

**Table 1. The genes and primers used for qRT-PCR experiments.**

| Gene   | Forward primer (5'-3') | Reverse primer (5'-3') |
|--------|------------------------|------------------------|
| MX1    | GACAGGACCATCGGAATCTTGAC | GGGCTTCGGACAGGCTCAG    |
| IFIH1  | GGTGGTGATGATGAGTATTGTGATG | AGATTATTCCTCCTGCTGATTCCCTC |
| IFIT1  | GTGGACCCTGAAAAACCCTGAATC | AGCGGACAGCCTGCCTTAG    |
| P4HB   | TGACGGCAAACTGAGCAACTTC  | TCGGTGTGGTGCTGCTGCG    |
| ISG15  | TGCTGGTGTTGGAACAAATGC   | CCCGCTCACTTGCTGCTTCC  |
| SOD2   | GCACCACAGCAAGCACCAC    | GATATGACCACCACCATTGAACCTTC |
| Actin  | TGGACTTCCAGCAAGAGATG   | GAAGGAAGGCTGGAAGAGTG   |

**Table 2. The all original data from the iTRAQ was analyzed using parameters.**

These data were uploaded in Table 2.

**Table 3. The 183 up-regulated DEPs were analyzed in OE VS NC.**

These data were uploaded in Table 3.

**Table 4. The 155 down-regulated DEPs were analyzed in OE VS NC.**

These data were uploaded in Table 4.

**Supplementary. The all original data included Table 2, Table 3 and Table 4.**

These data were uploaded in Supplementary.
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