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ORIGINAL RESEARCH

Promotion of miR-221-5p on the Sensitivity of Gastric Cancer Cells to Cisplatin and Its Effects on Cell Proliferation and Apoptosis by Regulating DDRI

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cancer cells to cisplatin, and the proregulating DDR1.

Patients and Methods: Altogether 69 patient, who treated with radical gastrectomy from January 2014 to January 2016 were collected. What the agree of the patients, 69 gastric carcinoma and 69 adjacenteessues were teen, respectively, during the operation, and gastric carcinoma and human gasting mucosa cell were purchased. RT-PCR was used for detection of the expression level of mn 221-5p d DDR1. Wound healing assay and CCK-8 assay of the cell migration and viability. Western blot and double were used for e plos ene y recommend to determine the target gene of miR-221-5p. luciferase reporter

a that miR-221-5p expression was decreased in GC tissues and cell Result vas sho The hi line expression of miR-221-5p reduced the resistance of GC cells to cisplatin and ibited th iferation and migration of gastric cancer cells. The high expression of miR-221 omoted the proliferation, invasion and migration of GC cells. In addition, we found that DD was a direct target gene of miR-221-5p in GC cells. We found that DDR1 expression creased in gastric carcinoma. Moreover, there was a negative correlation of R1 with the expression level of miR-221-5p. The increase of miR-221-5p increased the cheresensitivity of GC cells to cisplatin, and inhibited the proliferation, invasion, migration and EMT of GC cells by targeting DDR1.

Conclusion: The above research indicated that miR-221-5p may be a target for enhancing cisplatin chemotherapy sensitivity in gastric cancer patients.

Keywords: miR-221-5p, DDR1, gastric cancer cells, cisplatin, sensitivity, proliferation, apoptosis

Introduction

In recent years, gastric cancer has gradually become one of the main causes of death.¹ However, due to the delitescent early symptoms of gastric carcinoma, many patients are already at the advanced stage for first visit, and the prognosis of patients is often poor.^{2,3} In addition, chemotherapy resistance is one of the causes for the poor survival rate of gastric carcinoma patients.⁴ Cisplatin (DDP) is a firstline chemotherapy drug widely used in clinic at present, and is also widely applied for the treatment of gastric carcinoma. However, most patients will develop DDP

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resistance after a period of treatment, which is also one of the main reasons leading to recurrence or metastasis of patients.⁵ Therefore, how to improve the chemotherapy resistance of gastric carcinoma patients and the chemosensitivity of patients to DDP is one of the difficult problems to be solved clinically at present.

miRNA is a non-coding microRNA, which plays a very critical role in gene regulation and target protein degradation after transcription.⁶ In recent years, some researches have found that miRNA not only has a close relationship with the biological function of tumor cells, but also has certain influence on drug resistance of tumor cells.⁷ For example, previous studies⁸ found that miR-124 can inhibit cisplatin resistance of lung cancer cells by regulating STAT3. miR-221 is a miRNA closely relate to cancer, which can be processed into miR-221-3p or miR-221-5p to find the role of gene regulation.⁹ Different from miR-221-3p, the researches on miR-221-5p are rare in recent years, and only a few researches have reported the effect of miR-221-5p. For example, a study¹⁰ found that miR-221-5p acts as a tumor suppressor gene in prostatic carcinoma, but no research has been conducted on its role in gastric cancer and related mechanisms.

Disc domain receptor 1(DDR1), as a receptor tyrosine kinase, is up-regulated in various tumors and is latenced to regulate the biological function of tumor celt.⁴¹ In ocent years, studies¹² have found that the up-regulation control has a relationship with the poor processis of the patients, which leads us to suspect the there DDe1 has a relationship with the chemotherapy relistance of gastric carcinoma patients. However, no research has been conducted to discuss this.

Interestingly, throug o (http://www.targetscan.org/vert 72/), we found a constead a lationship between miR-221-5p and DDPr, but the mechanism of cisplatin resistance and biological function of cisplatin resistance tric carcinoma and s has not been studied. Therefore, we carried out this research in order to provide more molecular directions for the treatment of gastric carcinoma.

Materials and Methods

Clinical Specimen

Altogether 69 patients who treated with radical gastrectomy from January 2014 to January 2016 were collected. A total of 69 gastric carcinoma and 69 adjacent tissues were obtained with the consent of the patients and stored

Table I General Data of Patients

Data	Gastric Cancer Patients (n=69)
Sex	
Male	36(52.17)
Female	33(47.83)
Age (years)	62.15±9.26
BMI (kg/m ²)	22.25±1.06
Pathological Type	
Adenocarcinoma	25(36.23)
Squamous cell carcinoma	27(39.13)
Adenosquamous carcinoma	17(24.64)
Pathological Staging	
1	21(30.45
П	26(37.68)
ш	22(31, 3)
Degree of Differentiation	
High	(28.99)
Middle	23
Poorly	26(37.68)

in liquid nitrogen tanks. See Table 1 for patient information. Lee Table 2 or inclusion and exclusion criteria.

initional Review Board Statement

Tus study was reviewed and approved by the SIR RUN RUN Hospital, Nanjing Medical University Ethics committee.

Experimental Reagents and Materials

Human gastric carcinoma cell lines SUN-1, MKN-7, MGC-823, SGC-7901 and human normal gastric mucosa cell line GES (ATCC subordinate agent BeNa Biology, Beijing, China); Cisplatin (Shanghai Gold Wheat Biotechnology Co., Ltd.); qRT-PCR and reverse transcription kit

Table 2	2	Inclusion	and	Exclusion	Criteria
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Inclusion Criteria	Exclusion Criteria
Patients with gastric cancer diagnosed by pathology Patients with gastric cancer diagnosed for the first time	Patients who have received radiotherapy and chemotherapy Patients with other malignant tumors Patients with severe liver and kidney dysfunction Patients with severe infectious diseases Patients who refused to provide experimental specimens

(TransGen Biotech, Beijing, China); CCK-8 kit (Promega Company, USA); Transwell kit (Shanghai Fanke Biotechnology Co., Ltd.); PBS, fetal bovine serum (FBS) (Gibco Company, USA); Trizol reagent (Beijing Biolab Technology Co., Ltd.); double luciferase reporter gene detection kit (Beijing Biolab Technology Co., Ltd., KFS303-TFX); RIPA, BCA protein kit (Thermo Scientific Company, USA); Annexin V-FITC/PI apoptosis kit (Beijing Jiamay Biotech Co., Ltd., LHK601-020); DDR1, Caspase-3, Bax, Bcl-2 and β -Actin antibodies (Cell Signaling Technology Co., ltd.); goat anti-rabbit IgG secondary antibody (Boster Biotech, Wuhan, China); ECL developer (Thermo Co., ltd.); PCR instrument (ABI Co., ltd., USA). All primers were devised and compounded by Sangon Biotech Co., Ltd, Shanghai, China.

Cell Culture and Transfection

Human gastric carcinoma cell lines SUN-1, MKN-7, MGC-823, SGC-7901 and human normal gastric mucosa cell line GES were placed in DMEM medium including 10% PBS and 100U/mL of penicillin and streptomycin, and cultured at 37°C with 5% of CO₂. When the adherence growth and fusion of cells were observed to reach 85%, 25% pancreatin was added for digestion. After dig the cells were placed in the medium for continuous curve and passage, and then the expression of miP-221-5p DDR1 in each cell line was detected. The, MG 823 a. SGC-7901 were selected for transferron, and miP-221 5p-inhibitor (suppression sequent), m. 7-5p-mimics (over-expression sequence), r negative ntrol (miR-NC), targeting inhibition of DDR RNA (si-DDR1), targeting over-expression f DDR1 R (sh-DDR1), and negative control RM, (NC) were transfected into MGC-823 and SGC-790 with pofectamine[™] 2000 kit, and the operation tops we strictly accordance with the kit instruction

Real-The Quantitative PCR

Altogether 3×6^6 cells and 100mg of tissue were taken and ground, then the otal RNA in tissues and cells was extracted by Trizol reagent, 5µg of total RNA were collected to reverse transcript of cDNA referring to the instructions, and 1µL of synthesized cDNA was taken for amplification after transcription. The amplification system was as follows: 1 µ l of cDNA, 0.4µL of upstream and downstream primers with the concentration of 2umol/L, 10 µ l of TransScript[®] Tip Green qPCR SuperMix (2X), 0.4µL of Passive Reference Dye (50X), Nuclease-free Water was added to make up to 20µL.

U6 was taken as the internal reference of miR-221-5p, β -Actin as the internal reference of DDR1, and $2^{-\Delta \text{ CT}}$ was used to analyze the data.

Western Blot Test

Cell lysis and total protein extraction were carried out by RIPA lysis. BCA method was utilized to detect the protein concentration, and then the concentration was adjusted to 4µg/µL, 12% SDS-page electrophoresis separation was performed. After ionization, the sample was transferred to PVDF, and then 5% defatted mill powder was used to seal the PVDF membrane for firs. The DDR1 (1:500), Caspase-3(1:500), Bax (1:50, Bcl-2 (1:50), N-cadherin (1:500), E-Cadherin (1:500), imentin (1:500), and β-Actin (1:1000) primary antibody readded and sealed overnight at 4°C. fter shing with PBST to remove unbound printing anti-dies, the ARP-labeled goat antimouse sectory antibody (1, 000) was transferred to the membrane, and en incubated at 37°C for 1h. After that, the brane was vashed for 3 times with PBS, 5min/ me, and then illuminated with ECL and developed.

ell Proferation Test

The procession ability of MGC-823 and SGC-7901 cells evaluated by CCK-8 kit. Cells 48 hrs after transfection were collected, diluted to 3×10^4 cell/mL, and inoculated into 96-well plates. Each well was inoculated with 100μ L of cells, and cultured in 37° C with 5% CO₂. A 10μ L CCK8 solution was added to each well at 0h, 24h, 48h and 72h after the cells adhered to the wall. After the reagent was added, the cells were continuously cultured in an incubator at 37° C with 5% CO₂ for 2 h. Then, the OD value was measured at 450nm using an enzyme reader to detect the cell proliferation and visualize the growth curve. The experiment was repeated 3 times.

Apoptosis Test

Transfected cells were digested with 0.25% trypsin, washed twice with PBS after digestion, added with 100μ L of binding buffer, prepared into $1*10^6$ /mL suspension, sequentially added with AnnexinV-FITC and PI, placed at room temperature in dark for 5min, and detected with FACSVerse flow cytometer system. The experiment was repeated for 3 times to get the average value.

Cell Migration and Invasion Test

Cell migration and invasion were evaluated by scratchhealing test and Transwell test. For wound healing determination, 200µL aseptic pipette was used to scratch the cells to get a cell-free area, PBS was used to rinse the cells, and a new culture medium was added for culture. At 0h (W0) and 24h (W24) after cell scratch, the cell migration ability was evaluated by microscope for scratches at three different positions. Transwell assay: firstly, 200µL of DMEM culture solution containing 1x10⁵ cells was added into the upper chamber, and 500mLof DMEM containing 20% FBS was added into the lower chamber. Matrix and cells of the upper ventricle not passing through the membrane surface were cleaned after 48 hrs of culture at 37°C, rinsed with PBS for 3 times, fixed with paraformaldehyde for 10min, rinsed with double distilled water for 3 times, stained with 0.1% crystal violet for 10min after it was dried, and cell invasion was observed with a microscope.

Double Luciferase Assay

DDR1-3'UTR wild type (Wt), DDR1 1-3' UTR Mutant (Mut), miR-221-5p-mimics and miR-NC were transferred into SGC-7901 and SGC-7901/DDP cells by LipofectamineTM 2000 kit, and luciferase activity was detected by double luciferase reporter gene assay (Promega) 48 hrs after transfection.

Evaluation of Cisplatin Sensitivity

Cells were inoculated into 96-well plates with 1x10⁵ cells/ well. A 2mL of DDP with concentrations g 0.01µ mL, 2.5µg/mL, 5µg/mL, and 10µg/mL were added respe 11011 After incubation for 48h, fresh cutare am was was tran replaced, and 10µL CCK-8 solut cted to each well. Then, the cells were place into an incubator to continue culturing for 2b After that, a rbance values of each well were meaned at A50nm wavelength using SpectraMax M5 microster rester to detect cell proliferation. The experiment was related for 5 times. Then IC50 of DDP was alcula d according to cell survival rate. meentration of 5µg/mL was selected Finally, DK with a to intervene the Is and the apoptosis rate was detected.

Statistical Method

In this study, SPSS20.0 was utilized to statistically analyze the collected data. GraphPad 7 software package was used to visualize the required pictures. Independent t test was adopted for inter-group comparison, one-way ANOVA for multi-group comparison, LSD-t test for post-event pairwise comparison, repeated measurement ANOVA for multi-time point expression. Bonferroni and Pearson test were used for back testing to find out the correlation between miR-221-5p and DDR1 in the tissue. A P value less than 0.05 was considered a statistical difference.

Results

Expression Level and Clinical Meaning of miR-221-5p and DDR1 in Gastric Cancer RT-PCR detection results showed that compared with miR-221-5p in paracancerous tissues (1.07± 0.02), miR-221-5p in gastric cancer tissues was significantly decreased (0.42 ± 0.08) (P< 0.05), and compared with the expression of DDR1 in paracanceror assure (1.01 ± 0.12) , the expression level of DDR1 in stric cancer ssues was significantly increased $(1.84\pm0.21) \ge 0.05)$. The expression of miR-221-5p and LOR1 was in ative correlated (Figure r= -0.667, P< 05). Af , analyzing miR-221-5p, DDR1 and clinic athou feature, we found that miR-221-5p ar DDR1 has a se relationship with tumor diffe ntiak , TNM siging, and lymph node metastasis (P < 0.05). tients were divided into high and pression groups according to the average expression low iR-221-5p ┪ tumor tissues, with 36 cases in high of sion group and 33 cases in low expression group. expi Kaplan Jejer avival curve showed that the overall surrate of patients in high expression group was vious, higher than that in low expression group. Then, Cox regression analysis was carried out and it was oncluded that the expression of miR-221-5p was an independent risk factor for poor prognosis of gastric carcinoma, as shown in Figure 1, Tables 3 and 4.

Role of miR-221-5p on Cell Proliferation, Invasion, Migration, and Apoptosis

By detecting the expression of miR-221-5p in SUN-1, MKN-7, MGC-823, SGC-7901, and normal gastric mucosa cell line GES, we found that the expression of miR-221-5p in gastric cancer cells SUN-1, MKN-7, MGC-823, SGC-7901 was significantly lower than that in GES cells. Compared with the cells transfected with miR-NC, the expression of miR-221-5p in cells transfected with miR-221-5p-mimics by MGC-823 and SGC-7901 was obviously increased, and the expression transfected with miR-221-5p-inhibitor was obviously decreased. Detection of cell biological functions of the two groups showed that the proliferation, invasion and migration ability of transfected miR-221-5pmimics cells were significantly decreased. The proliferation, invasion and migration ability

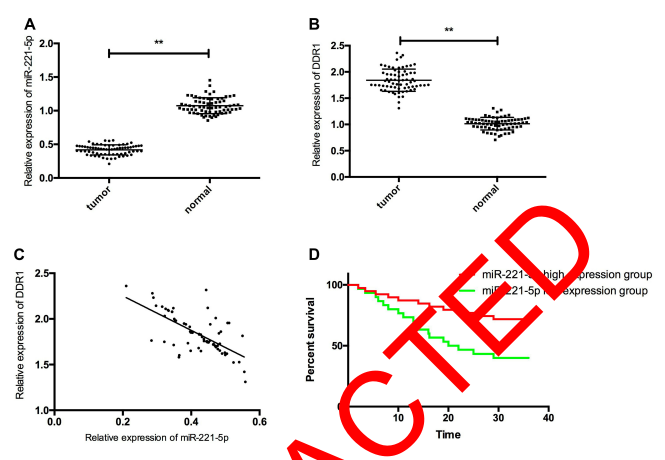


Figure I Expression and clinical significance of miR-221-5p and DDRI in patric case pression of miR-221-5p in gastric cancer tissue; (\mathbf{B}) expression of DDRI in gastric cancer tissue; (\mathbf{C}) miR-221-5p and DDRI were negatively correlated a gast patricer tissue; (\mathbf{D}) the overall survival rate of patients with miR-221-5p high expression group was significantly higher than that of patients miR-221-3p miR-221-3p and patients miR-221-3p miR-2

of transfected miR-221-5p-inhibitor lls cantly increased, and the apopted significantly s rate iR-330-3p decreased. After transfecting imics, the expression level of N-cadherin, pentin and Bcl-2 in aced, E-Cadhe cells was obviously r . Caspase-3 and Bax proteins were egnificately increased, the expression a Bcl-2 in transfected miR-221of N-cadherin, vime in 🖊 vignific Atly increased, and the 5p-inhibitor was n, Caspase-3 and Bax proexpressi level E-Cao teins w. luced. Figure 2. signi

Over-Expression of miR-221-5p on Cisplatin Sensitivity of Cells

After treating MGC-823 and SGC-7901 cells with different concentrations of cisplatin for 48h, CCK-8 results showed that under the intervention of cisplatin, compared with miR-NC group, the survival rate in miR-221-5pmimics group was decreased, and the survival rate in miR-221-5p-inhibitor group was increased, and showed DDP concentration dependent. We found through calculation that the IC50 of cisplatin in miR-221-5pmimics group cells was significantly reduced compared with miR-NC group, while that in miR-221-5p-inhibitor group was significantly increased. Finally, we selected DDP with a concentration of 5μ g/mL to analyze cisplatininduced apoptosis, and found that over-expression of miR-221-5p accelerated cisplatin-induced apoptosis, and the expression level of Caspase-3 and Bax proteins in cells was obviously increased, and the expression level of Bcl-2 protein was significantly reduced. Figure 3.

Role of DDRI Expression on Cell Proliferation, Invasion, Migration and Apoptosis

By detecting the expression level of DDR1 in SUN-1, MKN-7, MGC-823, SGC-7901 and human normal gastric mucosa cell line GES, it was showed that the expression of DDR1 in gastric carcinoma cells SUN-1, MKN-7, MGC-823, SGC-7901 was significantly increased. MGC-823 and SGC-7901 transfected cells with Si-DDR1 significantly

Factor		miR-221-5p Relative Expression	T value	P value	DDR1 Rela Expression		T valu	e P value
Sex	Male (n=36) Female (n=33)	0.42±0.07 0.41±0.08	0.554	0.582	1.85±0.21 1.84±0.21		0.198	0.844
Age	<62 years old (n=32) ≥62 years old (n=37)	0.43±0.08 0.41±0.07	1.108	0.272	I.86±0.20 I.83±0.22		0.589	0.558
TNM Staging	I, II (n=47) Illa (n=22)	0.46±0.05 0.33±0.04	10.69	<0.001	1.72±0.13 2.09±0.12		11.28	<0.001
Pathological Type	Adenocarcinoma (n=25) Squamous cell carcinoma (n=27) Adenosquamous carcinoma (n=17)	0.41±0.09 0.42±0.06 0.44±0.07	0.827	0.442	1.87±013 1.8100.18 1.80⊾		0.538	0.586
Lymph Node Metastasis	Not transferred (n=40) Transferred (n=29)	0.47±0.04 0.34±0.04	14.44	-0.001	1.70.0.14 2.04±0.14		10.79	<0.001
Degree of Differentiation	Low differentiation (n=26) Medium and high differentiation (n=43)	0.35±0.06 0.46±0.05	8	<0.00	2.06±0.14 1.71±0.12		11.02	<0.001
Table 4 Cox Ana Variable	alysis	Y	Uni	variate Ar	nalvsis	Multiv	ariate A	nalvsis
			P	HR	95% CI	P	HR	95% CI
Sex (male vs female) Age (<62years vs ≥62 yea Pathological types (adenocarc van, phosphores cancer vs adenosquamous carcinoma)			0.38 0.45 0.37	5 0.752	0.339–1.511 0.361–1.533 0.354–1.512			
Pathological serve (I+II strue vs III structure) Lymph node matastasis (III structure) Degree of difference don (low vs medium+high) miR-204(High vs Lou			0.02 0.00 0.03 0.00	3 2.891 2 1.973	1.314-4.485 1.372-4.793 1.092-3.576 1.592-8.216	0.032 0.009 0.602 0.006	2.916 2.455 1.069 3.362	1.083–7.886 1.296–4.122 0.814–4.019 1.304–4.126

Table 3 Relationship of miR-221-5p, DDR1 with Pathological Data of Patients

decreased the expression of DDR1 compared with cells transfected with Si-NC. Examining the biological functions of the two groups of cells, we found that the expression of DDR1 in cells transfected with Si-DDR1 by MGC-823 and SGC-7901 was obviously decreased than that in cells transfected with Si-NC, and the expression of DDR1 in cells transfected with Sh-DDR1 was significantly higher.

Detection of cell biological functions of the two groups showed that the proliferation, invasion and migration of cells transfected with Si-DDR1 were obviously decreased, and the apoptosis rate was significantly increased. The proliferation, invasion and migration ability of transfected Sh-DDR1 cells were significantly increased, and the apoptosis rate was markedly decreased. Compared with Si-NC group,

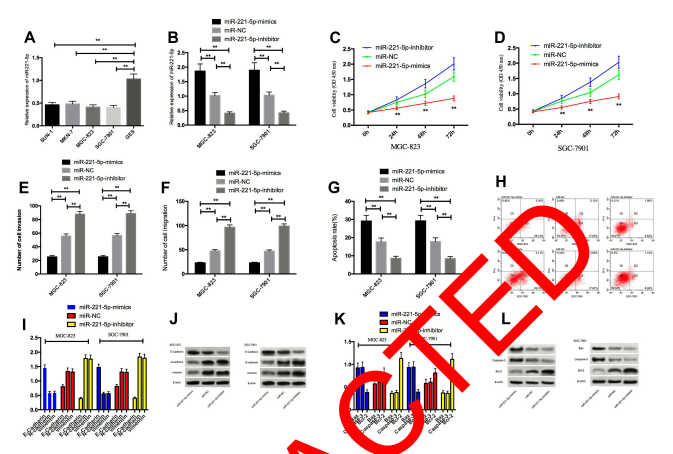


Figure 2 Effect of miR-221-5p on cell proliferation, invasion, migration and miR-221-5p low expression in gastric cancer cells; (**B**) expression of miR-221-5p DOD after transfection; (C and D) effects of up-regulating miR-221-5p and dow -5p expression on proliferation of gastric cancer cells; (E) effect of upregula tric cancer cells; (**F**) effect of up-regulating miR-221-5p and down-regulating miR-221regulating miR-221-5p and down-regulating miR-221-5p expression on invasio ing miR 5p expression on gastric cancer cell migration; (G) effect of -5p and down-regulating miR-221-5p expression on apoptosis of gastric cancer cells; (H) niR-221 Flow cytometry of apoptosis; (I) effects of up-regulating regulating miR-221-5p expression on EMT-related proteins N-cadherin, vimentin and and do E-Cadherin in gastric cancer cells; (J) protein image; (effect of u egulating m 21-5p and down-regulating miR-221-5p expression on apoptosis-related proteins Bax, Bcl-2 and Caspase-3 in gastric cancer cells; (L) protei 0.05 ge.

the expression of N-cadherin, vince in and Bch. in transfected Si-DDR1 cells as significant reduced, and the nerin, Caspase-3 and Bax proteins expression of E-C was markedly inc. sed the expression of N-cadherin, vimentin and 1.2 in unsfecte Sh-DDR1 cells was sigexpression of E-Cadherin, nificantly increas l, and and and steins was significantly reduced. Caspas Figure 4.

Inhibition of DDR1 Expression on Cisplatin Sensitivity of Cells

After treating MGC-823 and SGC-7901 cells with different concentrations of cisplatin for 48h, CCK-8 results showed that under the intervention of cisplatin, compared with Si-NC group, the survival rate of cells in Si-DDR1 group was lower, and the survival rate of cells in Sh-DDR1 group was higher, and showed DDP concentration dependent. Through calculation, we found that compared with Si-NC group, the IC50 of cisplatin in cells in Si-DDR1 group was significantly reduced, the cisplatin IC50 in Sh-DDR1 cells increased significantly. We selected DDP with a concentration of 5μ g/mL to analyze cisplatin-induced apoptosis. It was found that inhibiting DDR1 expression enhanced cisplatin-induced apoptosis, and the expressions of Caspase-3 and Bax proteins in cells increased markedly, while the expression level of Bcl-2 protein decreased markedly. Figure 5.

Identification of miR-221-5p Target Gene

For further verifying the correlation of miR-221-5p with DDR1, firstly, a target binding site was found between DDR1 and miR-221-5p by predicting miR-221-5p downstream target genes through Targetscan7.2. For this reason, we carried out double luciferase activity detection. The results showed that pmirGLO-DDR1-3'UT Wt luciferase

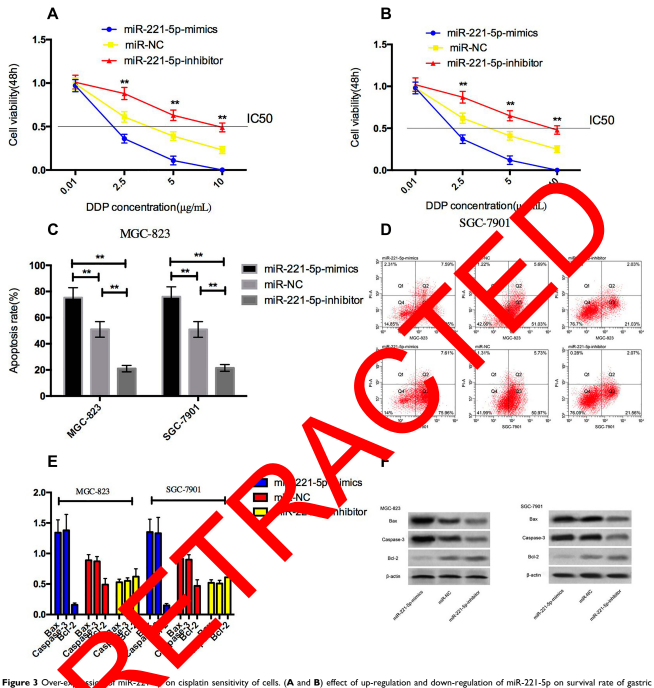


Figure 3 Over-expression of miR-221-5p on cisplatin sensitivity of cells. (A and B) effect of up-regulation and down-regulation of miR-221-5p on survival rate of gastric cancer cells under cisplatin intervention; (C) effect of up-regulation and down-regulation of miR-221-5p on apoptosis rate of gastric cancer cells under cisplatin intervention; (D) Flow cytometry of the proteins in gastric cancer cells under cisplatin intervention; (E) effect of up-regulation and down-regulation of miR-221-5p on apoptosis-related proteins in gastric cancer cells under cisplatin intervention; (F) protein in. ** indicates that P<0.05.

activity was significantly reduced after miR-221-5p overexpression (P< 0.05), but it had no influence on pmirGLO-DDR1-3'UTR Mut luciferase activity (P> 0.05). WB detection found that the expression of DDR1 protein in MGC-823 and SGC-7901 cells was significantly reduced after transfecting miR-221-5p-mimics, and the expression of DDR1 protein in the transfected microRNA-221-5pinhibitor group was markedly increased (P < 0.05), as shown in Figure 6.

Discussion

Among malignant tumors, gastric cancer is a common digestive system tumor, and its main treatment methods are surgery and systemic chemotherapy.^{13,14} However,

that P<0.05.

Sh-DDR1

Si-NC

Si-DDR1

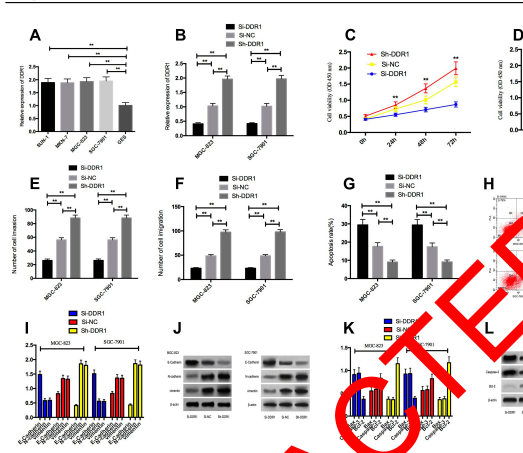


Figure 4 Effect of DDR1 on cell proliferation, invasion, migration and high expression of DDRI in gastric cancer cells; (B) expression of DDRI after Dto transfection; (C and D) effects of up-regulating DDRI and down-regulating proliferation of gastric cancer cells; (E) effect of up-regulating DDRI and DRI e down-regulating DDRI expression on invasion of gastric cancer cells; (F) up-regulating DDRI and down-regulating DDRI expression on gastric cancer cell migration; (G) effect of up-regulating DDRI and down-regulating l expr on on apoptosis of gastric cancer cells; (H) Flow cytometry of apoptosis; (I) effect of up-*L*MT-rela regulating DDR1 and down-regulating DDR1 expression cadherin, vimentin and E-Cadherin in gastric cancer cells; (J) protein image; (K) effect of l protein up-regulating DDRI and down-regulating DDRI exp on on app osis-related pteins Bax, Bcl-2 and Caspase-3 in gastric cancer cells; (L) protein image. ** indicates

since most patients were la stage whe they are diagnosed, they have lot the opportunity of surgery and th chemotherapy. Due to the exiscan only be treated tence of drug rest ance to chemotherapy, many patients ven ine ctive treatment, which will suffer from poor e.^{15,16} dise will furthe aggh ate th

, the effect of miRNA in tumor has also In 1 ent year been exp. red nore accery. In the process of tumor occurrence and de lopment, miRNA can regulate the biological function of the nor cells by regulating tumor suppressor genes or oncogenes.^{17,18} In our research, we also proved that miR-221-5p has a role of cancer suppressor in gastric cancer. The relationship between microRNA and chemotherapeutic drug resistance has attracted more and more attention. For example, microRNA-217 can improve the sensitivity of non-small cell lung cancer to cisplatin by targeting KRAS.¹⁹ Although the role of miR-221-5p in tumor has also been explored in relevant studies, the role of miR-221-5p in gastric cancer and its influence on cisplatin sensitivity have not been studied and discussed, so we have also explored it. This study found that the expression of miR-221-5p in gastric carcinoma was significantly decreased, and up-regulation of miR-221-5p could effectively restrain the proliferation, invasion and migration of gastric carcinoma and promote the apoptosis. For further understanding the role of miR-221-5p on cisplatin sensitivity, we regulated the expression of miR-221-5p in the case of cisplatin intervention, and observed that the increase of miR-221-5p improves the sensitivity of gastric cancer cells to cisplatin, while the down-regulation of miR-221-5p reduces the sensitivity of gastric cancer cells to cisplatin, which suggested that miR-221-5p may be a potential target for improving the chemotherapy efficacy of gastric cancer patients.

DDR1 is a member of transmembrane receptor tyrosine kinase, and many studies^{20,21} show that when DDR1

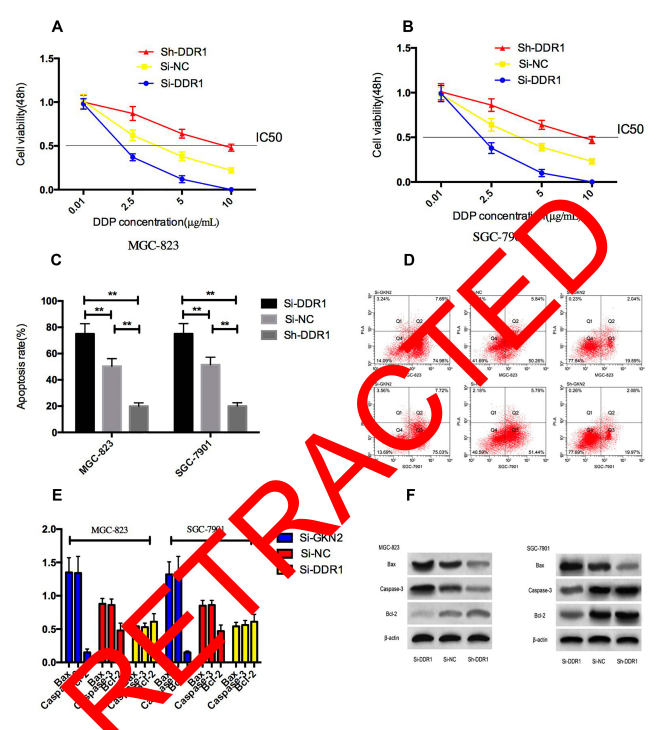


Figure 5 Inhibition of DDR pression on cisplatin sensitivity of cells. (A and B) effect of DDRI up-regulation and down-regulation on survival rate of gastric cancer cells under cisplatin intervention; (C) effect of DDRI up-regulation and down-regulation on apoptosis rate of gastric cancer cells under cisplatin intervention; (D) Flow cytometry of apoptosis; (E) effect of up-regulation and down-regulation of DDRI on apoptosis-related proteins in gastric cancer cells under cisplatin intervention; (F) protein image. ** indicates that P<0.05.

signaling pathway is activated by collagen, it can enhance the proliferation and migration of tumor cells, and DDR1 may be a cancer-promoting factor in cancer. Our research found that DDR1 is increased in gastric carcinoma, and silencing DDR1 expression can effectively inhibit the proliferation and invasion of gastric carcinoma and promote the apoptosis of gastric cancer cells. The result is consistent with previous studies.¹² Epithelial-interstitial transformation (EMT) is a process in which epithelial cells lose their original polarity, thus obtaining anti-apoptosis, high

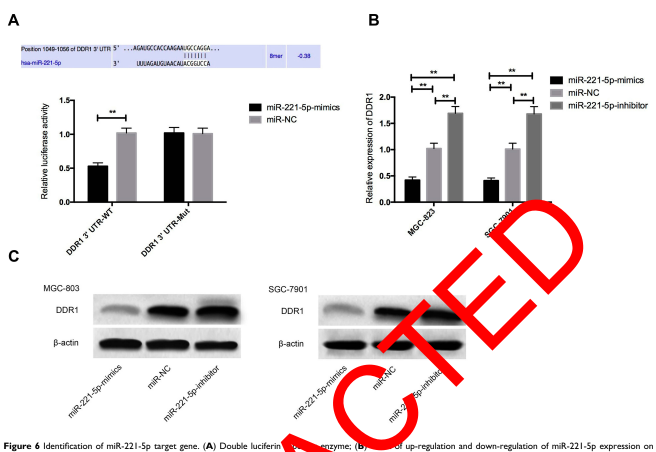


Figure 6 Identification of miR-221-5p target gene. (A) Double luciferin DDRI protein; (C) protein image. ** indicates that P<0.05.

migration, and invasion ability, and transforming to int stitial cells.²² However, the occurrence of DDR and EM is also closely related. Reports²³ e sh that the upregulation of DDR1 can prome the develop ent of colorectal cancer by reducing the expression of E-Camerin. We also found that silencing DR1 expression in gastric cancer cells can effectively accelerate the sensitivity of gastric carcinoma to cisple in, her ever, up-regulating the expression of DDR1 reduct the sense vity of nevertheless to cisplatin. s no **have** a previous research on the here \ sensitienty of D^V 21 to cisplatin, so we still do not know how DDL Lects the sensitivity of cisplatin, and further research is hedd. Vimentin, N-cadherin, E-cadherin, as markers for EM cransformation, their abnormal expression is also a feature of EMT.^{24,25} Previous studies^{26,27} have reported that abnormal activation of EMT will lead to the conversion of E-cadherin into N-cadherin protein, which will promote the formation of direct adhesion between cells and matrix. Vimentin, as an intermediate filament of matrix, will also lead to the adhesion and metastasis of cells, which is the key to abnormal activation of EMT. And we found that its mechanism may be realized by

regulating DDR1 protein. Previous studies²⁸ have also found that DDR1 can enhance invasion, metastasis and EMT of gastric cancer cells, and²⁹ have found that inhibition of DDR1 can prevent peritoneal metastasis of gastric cancer. All above results have confirmed our conclusions, but have not explained the upstream mechanism of DDR1. In this study, it was showed that DDR1 and miR-221-5p have targeted sites through the detection of miR-221-5p target genes. Therefore, we found that miR-221-5p can target and adjust the expression of DDR1 through double luciferase report detection, and the expression of DDR1 is significantly reduced after over-expression of miR-221-5p, which indicates that miR-204 can target and adjust the expression of DDR1 to restrain the proliferation, invasion, migration and EMT progress of gastric carcinoma, promote the apoptosis of gastric carcinoma and improve the sensitivity of gastric cancer cells to cisplatin.

To sum up, miR-221-5p over-expression gastric cancer patients have poor prognosis. miR-221-5p over-expression can inhibit proliferation, invasion, metastasis and EMT of gastric carcinoma by mediating DDR1, promote apoptosis and improve cisplatin sensitivity. However, there are still

some deficiencies in this study. First of all, we have not carried out nude mouse tumorigenesis experiment, so it is not clear whether miR-221-5p has influence on tumor size after cisplatin intervention in nude mouse. Secondly, this study has not detected the DDR1 signal pathway, only detected the expression of DDR1, and the specific mechanism of action is needed to be further verified. Therefore, we hope to carry out more basic experiments in future research to address our research deficiencies.

Conclusion

This study initially proved that miR-221-5p is downregulated in gastric carcinoma and the low expression of miR-221-5p has a relationship with poor prognosis of gastric cancer patients. Moreover, up-regulation of miR-221-5p can also inhibit the proliferation, invasion, migration and EMT of gastric carcinoma, promote the apoptosis of gastric carcinoma, and improve the sensitivity of gastric cancer cells to cisplatin. These findings can provide an important target direction for gastric carcinoma patients to better improve the efficacy of cisplatin chemotherapy.

Disclosure

The authors report no conflicts of interest in this work.

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