#### 8 Open Access Full Text Article

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## ORIGINAL RESEARCH **RETRACTED ARTICLE: SRF Potentiates Colon** Cancer Metastasis and Progression in a microRNA-214/PTK6-Dependent Manner

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Tao Li<sup>1,\*</sup> Yingchun Wan<sup>2,\*</sup> Ziyuan Su<sup>3</sup> Jiayu Li<sup>4</sup> Minna Han<sup>5</sup> Changyu Zhou D<sup>6</sup>

<sup>1</sup>Department of Anesthesiology, China-Japan Union Hospital of Jilin University, Changchun 130033, Jilin, People's Republic of China; <sup>2</sup>Department of Endocrinology, China-Japan Union Hospital of Jilin University, Changchun 130033, Jilin, People's Republic of China; <sup>3</sup>Department of Pharmacy, Changchun Second Hospital, Changchun 130062, Jilin, People's Republic of China; <sup>4</sup>Department of Cardiology, China-Japan Union Hospital of Jilin University, Changchun 130033, Jilin, People's Republic of China; <sup>5</sup>Department of Medicine, Medical School of Ching People & Apos's Liberation Army, Chinese People & Apos's Lik ation Army General Hospital, Poing 100853, People's Republic of Chia; <sup>6</sup>Departmen hent of Digestion, China-Japa ospital Unio 30033, of Jilin University Changer of Ch Jilin, People's

outed equally to \*These nors con this work

Correspondence: Changyu Zhou Department of Digestion, China-Japan Union Hospital of Iilin University. Changchun 130033, Jilin, People's Republic of China Tel/Fax +86-431-84995850 Email drZhouCY1224@163.com



e-specific trans Objective: Serum response factor (SRF), a seque int related to metastasis of gastric cancer, a digestive tract concer. Herein, we probed the effect C), another digestive tract disorder, of SRF on metastasis and progression of certain can and the detailed mechanism. Methods: Microarray analysis we condited on turn, and adjacent tissues to filter

differentially expressed miRNA, followed by T-qPCR validation in CC cell lines. The transcription factor and the reget gene of microl A-214 (miR-214) were predicted, and their binding relationship were tested y luciferase reporter assays and ChIP assays. ein tyrosine nase 6 (PTK6) expression in CC patients and Subsequently, SRF and pr while J K2 and STAT3 expression in cells by Western blot cells was evaluated by RT-qP analysis. To furt lore functions of miR-214, PTK6 and SRF on CC, CC cells were mimic and/or SRF overexpression. delivered with si-K6, r

ssed poorly in CC tissues and cell lines, which related to advanced Result R-214 vival. miR-214 mimic inhibited proliferation, migration, invasion, TN stagin and st for grow and metastasis of CC cells. SRF, overexpressed in CC samples ograft 🕇 JV suppressed the transcription of miR-214. Meanwhile, SRF upregulation counterand acted the inhibitory role of miR-214 mimic in CC cell growth. miR-214 negatively regulated PTK6 exp. sion to impair the JAK2/STAT3 pathway activation, thereby halting CC cell liferation, migration, invasion, xenograft tumor growth and metastasis.

Collusion: Altogether, miR-214 may perform as a tumor suppressor in CC, and the SRF/ miR-214/PTK6/JAK2/STAT3 axis could be applied as a biomarker and potential therapeutic target.

Keywords: SRF, microRNA-214, PTK6, Colon cancer, The JAK2/STAT3 pathway

### Introduction

As for the cause of cancer-related death, the ranking of colon cancer (CC), also named as colorectal cancer, showed a downward trend (the third place) in the United States by 2014, partly as a consequence of historical changes in risk factors, the use and spread of early detection tests, as well as improvements in therapeutic interventions.<sup>1</sup> Nevertheless, incidence is still higher in men than women and strongly associated with age, and the median age at diagnosis is around 70 years old in developed countries.<sup>2</sup> Even though surgery can cure patients with CC at early stage, the 5-year survival rate is dismal for patients with metastatic CC (only 10%).<sup>3</sup> Metastasis is a complicated event that involves a series of procedures

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where tumor cells proliferate, detach from the primary tumor sites, and migrate to a distant organ.<sup>4</sup> Therefore, clarification of the mechanisms underlying the metastasis of CC may be beneficial to the discovery for new diagnostic biomarkers and effective treatment methods.<sup>5</sup>

MicroRNAs (miRNAs) are short non-coding RNAs with 19-25 nucleotides in length that function importantly, practically in cancer-related processes involving proliferation, cell cycle control, apoptosis in addition to migration.<sup>6–8</sup> Among them, miR-214 has been reported to play a tumor suppressor role in multiple cancers, including gastric cancer,9 pancreatic cancer,10 non-small lung cancer<sup>11</sup> as well as osteosarcoma.<sup>12</sup> Also, miR-214, notably reduced in CC, was modulated by a transcription factor FOXD3 and tightly correlated with lymphatic metastasis.<sup>13</sup> Also, the proliferation and invasion of CC SW620 cells were weakened by miR-214, yet the apoptosis rate was enhanced.<sup>14</sup> Moreover, serum response factor (SRF), another transcription factor regulating the expression pattern of over 200 genes, was observed to be hypermethylated in gastric carcinoma metastasis, signifying its function as an attractive biomarker for predicting gastric carcinoma metastasis and prognosis.<sup>15</sup> Therefore, we postulated that SRF may mediate the expression of miR-2 to influence the metastasis of CC. Protein tyrosine kinase (PTK6), an intracellular tyrosine kinase that preposed to facilitate apoptosis in CC,<sup>16</sup> has been validate as a rget gene of miR-214 in CC cells in the present w - In addition, endogenous PTK6 enhance cell val and oncogenic signaling following D damage CC.<sup>17</sup> Taken together, our study spectrated at miR-214 regulated the metastasis and progression of C by directly targeting 3'UTR of the K6 gene in an SKF-dependent manner. Gain- and loss f-fun on assays were carried out in CC LOVO and SW620 Is as we as in nude mice to verify our hy chesis

#### Materials and Methods

After the determinition of miR-214 using microarray analysis, RT-qPCR was carried out to assess its expression. For the purpose of examining the effects of SRF, miR-214 and PTK6 on CC, EdU, Transwell, tumor growth and metastasis assays were performed. Dual-luciferase reporter assays were used to detect targeting relationships between genes. Afterwards, ChIP was utilized to detect the binding relationship between SRF and miR-214. Western blot was finally applied to detect the activity of the JAK2/STAT3 signaling.

#### Clinical Data

From August 2014 to August 2016, 80 patients with CC (48 males and 32 females) treated in China-Japan Union Hospital of Jilin University were enrolled. The average age of the patients was  $59.06 \pm 9.08$  years. The clinical stage of patients was 17 in stage I, 28 in stage II, 23 in stage III and 12 in stage IV. Adjacent tissues were extract from 5-10 cm away from tumor tissue. The patients with radiotherapy, chemotherapy or any immunotherapy were excluded from this study. All patients with complete clinical er and kidnev data received blood routine, urine rout function, electrocardiogram and ther example nations to exclude the existence of chronic decises or con lications. All patient cases were ind pendently eviewed histologically as primary CC. The clinic pathole I features of the enrolled patients are more ized in Table 1. The present study was review a and approved by the Ethics Committee of China-Jap U. on Hospital Jilin University, and we obtained informed correct from all patients.

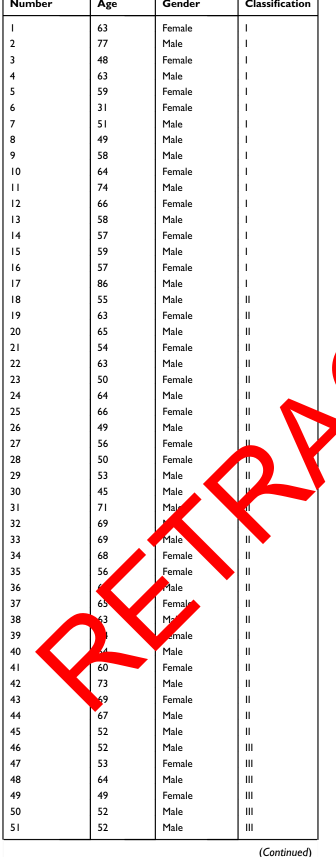
#### Mi roarray nalysis

tal miRN molecules of CC tissues and adjacent The ree patients were extracted using TRIzol tissues nt (Invitrogen, Carlsbad, CA, USA). RNA purity K. d concentration were determined by NanoDrop (Thermo Fisher Scientific Inc., Waltham, MA, USA). Following DNA synthesis using cDNA Reverse Transcription kits (Takara Biotechnology, Co., Ltd., Dalian, Liaoning, China), cDNA was labeled with a miRNA Complete Labeling and Hyb Kit (Agilent Technologies, Santa Clara, CA, USA). A total of 15 µg cDNAs was dissolved in 3 µL ddH<sub>2</sub>O and incubated with 1 µL calf intestinal phosphatase at 37°C for 30 min. The denatured RNA was inactivated at 95°C for 15 s and incubated with the mixture of 3 µL marker buffer, 1.5 µL Hy3 marker, 2 µL dimethylsulfoxide and 2 marker enzymes at 16°C in the dark for 1 h. Totally 125  $\mu$ L labeled RNA samples were incubated with 65  $\mu$ L hybridization buffer  $(2 \times)$  at 95°C for 2 min in the dark. The 200 µL mixture containing 15 µg cRNA was then hybridized with Human miRNA Microarray Release 14.0,8x 15K (Agilent) in a hybridization oven (hybridization oven) overnight, then scanned with Agilent SureScan Dx (Agilent). A quality control evaluation of the analysis results was carried out. Background correction and normalization of raw data was performed using Robust Multi-Array Average. miRNA was used analyzed and identified using *t*-test. miRNAs with p < 0.05 and |Fold change| > 1.5

 Table I Demographic Characteristics of the Enrolled Patients

 Number
 Age
 Gender
 Classification







was defined as differentially expressed miRNAs and plotted as a heat map by hierarchical clustering.

## RNA Isolation, cDNA Synthesis and Quantitative Polymerase Chain Reaction (qPCR)

The total RNA in tissues was extracted with the help of the TRIzol Reagent (Invitrogen). After the removal of the genomic DNA contamination, the template RNA was subjected to enzyme digestion. The purity and concentration of RNA was determined using a spectrophotometer. RNA integrity was detected by 1.5% agarose gel electrophoresis, and the RNA concentration was adjusted to  $500 \text{ ng/}\mu\text{L}$ . RNA samples were transcribed into cDNA using a cDNA Reverse Transcription kit (Takara Biotechnology). The SYBR RT-qPCR kit (Thermo Fisher Scientific) was used for amplification. Primers for this experiment were designed by Primer

Premier 5.0 (Premier Biosoft International, Palo Alto, CA, USA) and synthesized in Shanghai Sangon Biological Engineering Technology & Services Co., Ltd. (Shanghai, China). Glyceraldehyde 3-phosphatedehydrogenase (GAPDH) and U6 were treated as internal controls for SRF, PTK6 and miR-214, respectively. All primer sequences are listed in Table 2. The  $2^{-\Delta\Delta Ct}$  method was applied to measure the relative expression of mRNA and miRNA.

### Cell Culture and Treatment

CC cell lines SW620, SW480, LOVO, HCT 116, HT 29, LS174T, normal human colon fibroblast cells (CCD-18Co) and human embryonic kidney epithelial cells (HEK293T) were purchased from Keygen Biotech Co., Ltd. (Nanjing, Jiangsu, China). SW620 and SW480 were grown in a Leibovitz L-15 medium supplemented with 10% fetal bovine serum (FBS; Gibco, Grand Island, NY, USA) and LOVO, HCT 116, HT 29, LS174T and HEK293T cells were exposed to Roswell Park Memorial Institute 1640 (RPMI-1640) medium containing 10% FBS. All aforementioned cells were maintained in a 37°C cell incubator containing 5% CO<sub>2</sub>. Routine examination of mycoplasma in all cell lines showed negative results.

SRF fragments, miR-214 mimic and small interferi RNA (siRNA) targeting PTK6 (System Biosciences, Pal Alto, CA, USA) were cloned into a pmirG vector (Promega Corporation, Madison, WI, USA) and the constructed vectors were transfected into cells that stably overexpressing SRF, miR .6 deficit. .4 or 1 cells overe SRF-OE vectors were delivered ressing miR-214 in the rescue experiment. In offectamine 3000 (Invitrogen, Carlsbad, CA, SA) was appled for transient

#### Table 2 List of Primers Und in his Stud

Targets	Primer equent (5'-3
U6	Forward CTCGCTTCGGCAGCACA
GAPDH	Forvert: GGGAGCCAAAAGGGTCATCA Reverse, GATGGCATGGACTGTGGTC
SRF	Forward: TGCTGAATGCCTTCTCCA Reverse: GCCTGCTGCCCTATCACA
miR-214	Forward: CTGGGGGCTGGATTCGGGGGTGGG Reverse: ACCCCCACCCCGAATCCAGCCCCCA
РТК6	Forward: TACTTTGGGGAGGTCTTCGAG Reverse: TGCCGCAGCTTCTTCATG

Abbreviations: GAPDH, glyceraldehyde 3-phosphatedehydrogenase; SRF, serum response factor; miR-214, microRNA-214; PTK6, protein tyrosine kinase 6.

transfections of vectors and siRNA. Detection of transfection efficiency was carried out using RT-qPCR.

## 5-Ethynyl-2'-Deoxyuridine (EdU) Incorporation Assay

EdU assays were conducted to assess the viability of CC cells. LOVO and SW620 cells in good growth condition were seeded in a 24-well plate. The culture medium was added with the reagent from an EdU Staining Proliferation Kit (iFluor 488, Abcam Inc., Cambridge, UK) to a final concentration of 10 µmol/L, and incubated in the incubator The cells were then fixed with phosphate buffered ine (PBS) lution containing 4% paraformaldehyde at rochtemperature pr 15 min and incubated with PBS coptining 0. Triton-0 at room temperature for 20 min. A crwards the cent ach well were incubated for 30 min w 100 L Apollo 567 (Guangzhou RiboBio Co., Ltd. Guangz, 1, Guap Long, China) void of light and staine as 5 min with 4 amidino-2-phenylindole. Five visual nelds we randomly taken under a TCS SP8 roscope (L. a, Bannockburn, IL, USA). The confor luorescence indicates all cells, whereas the red fluoresblue is the repliciting cells infiltrated by EdU. The rate of cen EdUsitive cell was calculated.

#### In the well Assays

answell assays were carried out for cell migration and invation evaluation. Matrigel (BD Biosciences) was placed into the dical chamber under sterile conditions and cultured for 30 min. The apical chamber was then incubated with RPMI-1640 medium in a CO2 incubator for later use. After detachment and centrifugation, the cells were resuspended in serum-free medium and diluted into cell suspension ( $5 \times 10^5$  cell/mL). A total of 500 µL RPMI 1640 containing 10% FBS was supplemented to the basolateral chamber, and the diluted cell suspension (200 µL) was supplemented to the apical chamber for a 48-h culture in an incubator at 37°C with 5% CO2. Afterwards, cells that did not migrate or invade were removed by wiping them with a cotton swab. Cells left on the lower surface of the membrane were stained for 10 min with crystal violet. Five visual fields were arbitrarily chosen under a microscope (Leica DM500) to photograph the invading cells. Migration test was carried out without Matrigel precoating on the apical chamber, and the rest of the operation was consistent with invasion detection.<sup>18</sup>

### Tumor Growth Assay

Forty-eight specific-pathogen-free (SPF) female BALB/c nude mice (4–6 weeks old,  $20 \pm 2$  g) were from Beijing Vital River Laboratory Animal Technology Co., Ltd.

(Beijing, China). The nude mice were injected with LOVO or SW620 cells stably transfected with miR-214 mimic, miR-214 control, SRF-overexpression (OE), SRF-negative control (NC), miR-214 mimic + SRF-OE, miR-214 mimic + SRF-NC, si-PTK6 or PTK6-NC (n = 3). Totally  $4 \times 10^{6}$ LOVO and SW620 cells dispersed in 2 mL saline was injected subcutaneously into the nude mice. Mouse tumor volume was measured at an interval of 7 days after injection by formula:  $L \times W^2/2$  where L indicates length and W indicates width. After 28 days, mice were euthanized with 1% pentobarbital sodium at 120 mg/kg, and the tumor was weighed for histological experiments. All animal experiments were implemented as per the principles and procedures permitted by the Committee on the Ethics of Animal Experiments of China-Japan Union Hospital of Jilin University (Approval number: 20140216CJD).

#### Immunohistochemical Staining

The extracted mouse tumor tissues were embedded in paraffin, deparaffinized and hydrated. Five sections were obtained from each xenograft tumor. Afterwards, the sections were allowed to stand with 3% H<sub>2</sub>O<sub>2</sub> at room temperature for 15 min. After being blocked with normal goat serum for 15 min at room temperature, the sections were probed with primary antibody against KI67 (1:500, ab197234, Abca ) at 4°C overnight. Subsequently, the secondary artibody (1: ab199091, Abcam) was applied and incubied for 5 min ch perox 37°C. Following treatments with 40 the horser idase at 37°C for 15 min and diamin benz , the sections were counter-stained with hem explin for 30 dehydrated and sealed with neutral guy. Unc. a microscope (Leica DM500), five non-over pping fields the chosen for each section. The cells proventing brown or brown granules in the nucleus were define as K// positive.<sup>19</sup>

#### In vivo leta tasis says

Anothe 48 SPF come BALB/c nude mice (4–6 weeks old,  $20 \pm 2$  g) we used for in vivo metastasis assay. The nude mice were injusted with LOVO or SW620 cells stably transfected with mik 214 mimic, miR-214 control, SRF-OE, SRF-NC, miR-214 mimic + SRF-OE, miR-214 mimic + SRF-NC, si-PTK6 or PTK6-NC (n = 3). A total of  $4 \times 10^6$  LOVO and SW620 cells were inoculated into the nude mice through tail vein. After 45 days, the nude mice were euthanized with 1% pentobarbital sodium. The lung tissues of nude mice were paraffin-embedded and subjected to a hematoxylin-eosin (HE) staining (Beijing Solarbio Science & Technology Co., Ltd., Beijing, China) to observe

the formation of metastatic lesions. The sections were dewaxed with xylene for 8 min, soaked in gradient alcohol, immersed into hematoxylin for 15 min. The sections were differentiated in 1% hydrochloric acid alcohol for 30 s, immersed into eosin solution for 5 min, dehydrated with gradient alcohol, cleared for 10 min with xylene and sealed with gum. The lung sections were then visualized under a microscope (Leica DM500), and 5 fields were arbitrarily chosen for photographs.

#### Dual-Luciferase Assay

w.cuilab.cn TransmiR v2.0 (http://w ansmir) and ALGGEN (http://alggen.lsi.up s/) were i ed to predict the binding sequences etween S. and R-214 as well as between SRF and TK6 prinoter. Noinding sequences between miR-214 and PT to 3'untraslated region (3'UTR) were predict a by Stan se (bt .//starbase.sysu.EdU.cn/). PTK6 37 R suence and x-214 vector, PTK6 promoter sequence and SR vector, miR-214 promoter sequence and vector were instead into pMIR-REPORT<sup>TM</sup> (Thermo isher Scientific).<sup>20</sup> These vectors were co-transfected into EK293T cost using Lipofectamine 3000 (Invitrogen). 24 h de cells were lysed, and the luciferase activity intensity (intensity = RLU1/RLU2, RLU1 was firefly luciter, intensity, RLU2 was renilla luciferase intensity) was determined by a Dual-Luciferase Reporter Assay System (Promega Corporation).

## Chromatin Immunoprecipitation (ChIP) Assay

The ChIP was performed to validate the binding relationship between SRF and miR-214 using a Pierce Agarose ChIP Kit (Thermo Fisher Scientific). LOVO and SW620 cells were diluted with 37% formaldehyde to 1% concentration and incubated in a 37°C incubator for 10 min. After the addition of glycine, the cells at concentration of 0.125 M were allowed to stand at room temperature for 5 min. Following a 5-min centrifugation at 2000 rpm, the cells were lysed using sodium dodecyl sulfate (SDS) lysis buffer and proteinase inhibitor, and DNA was sheared by a sonicator (Shunma Tech., Nanjing, Jiangsu, China). After a 10,000 g centrifugation at 4°C for 10 min to remove insoluble substances, the samples were added with 900  $\mu$ L ChIP dilution buffer and 20  $\mu$ L 50  $\times$ protease inhibitor cocktail, and mixed with 60 µL ProteinA Agarose at 4°C for 1 h. Following a 10-min standing and a 700-rpm centrifugation, the supernatants were incubated with antibodies against SRF or IgG (ab2410, Abcam) at 4°C overnight. The immunoprecipitates were eluted using 250  $\mu$ L elution buffer, and the precipitated DNA was dissolved in 100  $\mu$ L ddH<sub>2</sub>O for the following PCR analysis.<sup>21</sup>

## Western Blot

The cells were lysed using radio immunoprecipitation assay buffer (Amresco, Radnor, PA, USA) and centrifuged at 800 g at 4°C for 5 min. Next, the cells were ice-bathed with 5  $\times$ lysis buffer for 10 min and centrifuged at 12,000 g for 10 min at 4°C to obtain the supernatant. Proteins were subjected to an SDS-polyacrylamide gel electrophoresis. Separated proteins were blotted onto polyvinylidene fluoride membranes (Millipore, Merck, Darmstadt, Germany) and sealed with 5% skim milk. The membranes were then immunoblotted with the specific primary antibodies against janus kinase 2 (JAK, 1:5000, ab108596, Abcam), p-JAK2 (1:2000, ab108596, Abcam), signal transducer and activator of transcription 3 (STAT3, 1:5000, ab68153, Abcam), p-STAT3 (1:1000, ab76315, Abcam) and  $\beta$ -actin (1:2000, ab32101, Abcam). After 16 h, the membranes were probed with the secondary antibody (1:3000, ab205718, Abcam). The optical density (OD) analysis of immunoblot images was performed using ImageJ software (version 1.8.0; National Institutes of Health, Bethesda, MD, USA).<sup>22</sup>

### Statistics

All the statistical analyses were implement a usii the SPSS 22.0 software (IBM Corp. Armonk, Y, US For all assays, at least 3 independent tria ried out. wer ndard devia Results are shown as the means  $\pm$ n (SD) and compared using a two-to-ted t- st for two-group comparison or one-way or wo-way analysis of variance (ANOVA) followed by Z key's pultiple comparison posthoc test for comparing fferer groups. The Kaplan-Meier and Log rank test were olied for evaluating survival b of less than 0.05 was comparisons. iled ptwo considered signif

## Results

miR-214 expression was poorly expressed in CC patients and cells which was not only negatively correlated with tumor, node, metastases (TNM) stage, but also showed good prognosis. The proliferation, invasion and migration of CC cells were weakened by miR-214 mimic. Also, the tumorigenic and metastatic potentials of CC cells were decreased in vivo after miR-214 overexpression. SRF bound to the miR-214 promoter and inhibited its transcription. SRF restored the activity of CC cells, while upregulation of SRF in cells overexpressing miR-214 also promoted the activity of CC cells. miR-214 targeted PTK6 and inhibited its expression, and depletion of PTK6 reduced the activity of CC cells. SRF/miR-214/PTK6 axis mediated the activity of the JAK2/STAT3 pathway, and SRF and PTK6 shared positive correlations with the pathway, while miR-214 had a negative correlation with the JAK2/STAT3 pathway.

## miR-214 is Significantly Reduced in CC Tissues and Cells

We first performed a miRNA mig array analy of tumor tissues and adjacent tissues in clinica. obtained ( patients. Among all differentially excessed mike As, y found that miR-214 was the most swnregy red one regure 1A). RTqPCR tests were performed tumor sues and adjacent tissues in all CC atients, which disp' yed that miR-214 was reduced in the row wes (Figure 3). We then analyzed the correlation between TNV staging and miR-214 expression in cents. It was demonstrated that miR-214 expression CC p negatively correlated with TNM staging in CC (Figure was 1C). Survival and vsis of CC patients displayed that high n of min-214 was more favorable for postoperative expres. ival (Figure 1D). Meanwhile, we examined the miR-214 ress. in LOVO, SW620, SW480, HCT116, HT-29, LS174T and CCD-18Co cells, and the results showed that iR-214 was also significantly lower in CC cell lines, and the downregulation was more pronounced in LOVO and SW620 cells (Figure 1E). LOVO and SW620 cells were then transfected with miR-214 mimic for subsequent experiments (Figure 1F).

## Overexpression of miR-214 Inhibits CC Cell Viability

We subsequently examined the involvement of miR-214 in cell growth. It was showed that the number of EdU-positive cells in cells overexpressing miR-214 was decreased significantly compared with the cell transfected with miR-214 control (Figure 2A). After 48 h, miR-214 mimic led to significantly reduced cell migration distance and invasive cell number (Figure 2B and C). Also, cells overexpressing miR-214 were subcutaneously injected into three nude mice, and the volume of subcutaneous tumor in mice with miR-214 mimic was reduced compared to those with miR-214 control (Figure 2D). The positive rate of surface marker KI67 was significantly decreased following miR-214 over-expression (Figure 2E). Even though mice in both groups

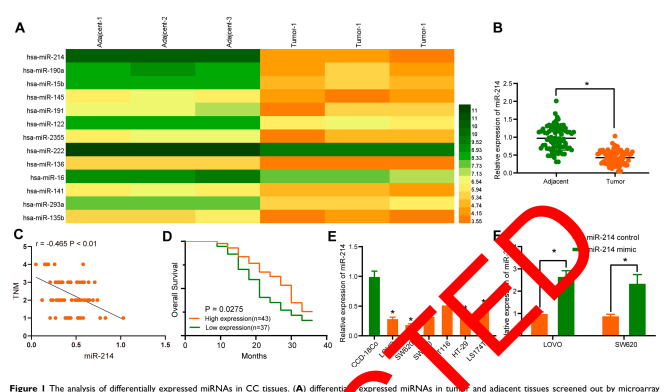


Figure 1 The analysis of differentially expressed miRNAs in CC tissues. (A) differential analysis; (B) miR-214 expression in CC tumor and adjacent tissues evaluated by RT-between miR-214 expression and TNM stage of patients with CC; (D) survival analysis (\*p < 0.05 according to the one-way ANOVA); (F) miR-214 mimic was transfected into

displayed metastasis dissemination. The pulmonary notules were notably diminished after overexpression of miR-24 and the area of individual pulmonary rodules was all significantly reduced (Figure 2F).

SRF Interacts with the NiR-214 omoter To examine the molecular mechanish of miR-214 in CC, we predicted the bip ing sites between the transcription factor SRF to its comote by TransmiR and ALGGEN (Figure 3A). The luc se report assays demonstrated that SRF rectually in pase the luciferase signals of miR-21 wild ty 2 (WT) (Ngure 3B). We then performed ChIP and sir to further test the combination of the two, and the result exhibited that SRF could directly bind to miR-214 promote region (Figure 3C). RT-qPCR detection of tumor tissues and adjacent tissues in CC patients illustrated that SRF expression in CC tissues was promoted (Figure 3D), which was also validated in CC cell lines as well (Figure 3E). These results indicated that SRF bound to specific promoters of miR-214 (Figure 3F). Therefore, LOVO and SW620 cells were infected with lentiviral vectors overexpressing SRF to construct SRF-OE cells for the subsequent experiments (Figure 3G).

#### CR (\*p < 0.05 according to the two-way ANOVA); (**C**) the correlation analysis f CC patients; (**E** uniR-214 expression in CC cell lines assessed RT-qPCR analysis OVO and SW6 cells (\*p < 0.05 according to the two-way ANOVA).

## Superior of SRF Enhances CC Cell Viability

The viability of SRF-OE cells was detected, and the number of EdU-positive cells was elevated, indicating the viability of cells was enhanced (Figure 4A). Meanwhile, the cell migration distance (Figure 4B) and the number of invaded cells (Figure 4C) were elevated significantly after 48 h. Moreover, the results of in vivo assays suggested that upregulation of SRF in three nude mice resulted in larger tumor volume (Figure 4D), higher KI67 positive rate (Figure 4E) as well as more and larger pulmonary nodules (Figure 4F) relative to the counterparts.

### SRF Inhibits the Transcription of miR-214

SRF-OE was then delivered into cells overexpressing miR-214 to construct miR-214 mimic + SRF-OE cells (Figure 5A). The viability of cells transfected with miR-214 mimic + SRF-OE was significantly increased, suggested by increased number of EdU-positive cells (Figure 5B). In support, the cell migration distance (Figure 5C) and the invasive cell number (Figure 5D) were upregulated after the administration of the miR-214 mimic + SRF-OE cell after 48 h. Three nude mice

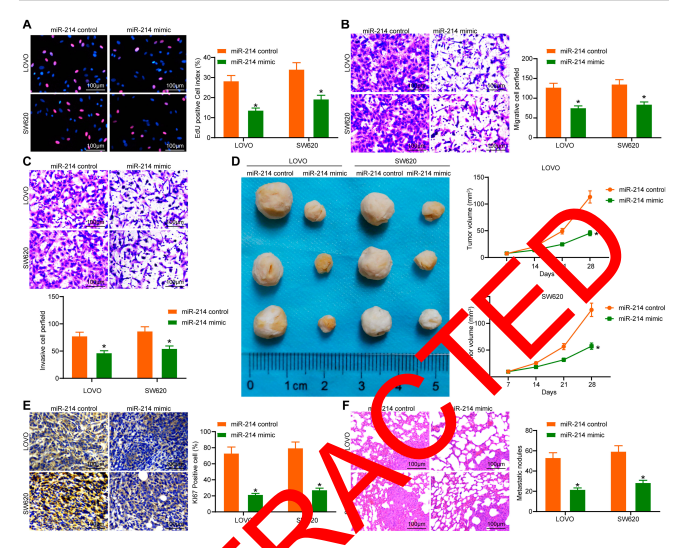


Figure 2 Increased miR-214 is associated with decreased CC or proliferative, migratory, invasive, tumorigenic and metastatic capacities. (A), CC cell proliferation evaluated by EdU staining; (B), CC cell migration of function ( $a_{12}, c_{12}, c$ 

that received miR-21 comining SRF-OE cell delivery appeared to exhibit a convier turnor burden (Figure 5E), greater K-67 politive site sigure 5F) and aggravated menstasis sigure 5G) versus nude mice received miK-12 mimic + SRF-NC cell delivery.

## miR-214 Negatively Regulates PTK6 Expression by Directly Binding

To identify downstream targets of miR-214, we used StarBase to predict PTK6 as a potential target. We cloned PTK6 3'UTR into a luciferase plasmid. In HEK293T cells, miR-214 mimic suppressed the luciferase activity of PTK6 3'UTR (Figure 6A). To determine whether PTK6 was regulated by SRF through a direct transcription, we overexpressed SRF to find out its impact on the dual luciferase activity of PTK6-WT and PTK6-MT. It turned out that SRF-OE had no notable effect on of the PTK6 promoter relative to the SRF-NC (Figure 6B). RT-qPCR detection of CC patients showed that PTK6 expression in CC tissues was significantly enhanced (Figure 6C). We identified a similar trend in CC cell line (Figure 6D). Moreover, the PTK6 expression decreased remarkably in the cells overexpressing miR-214 and increased significantly in the cells overexpressing SRF (Figure 6E). We may conclude that SRF modulated the PTK6 expression in a miR-214-dependent manner. As a result, cells with PTK6 knockdown were constructed for subsequent experiments (Figure 6F).

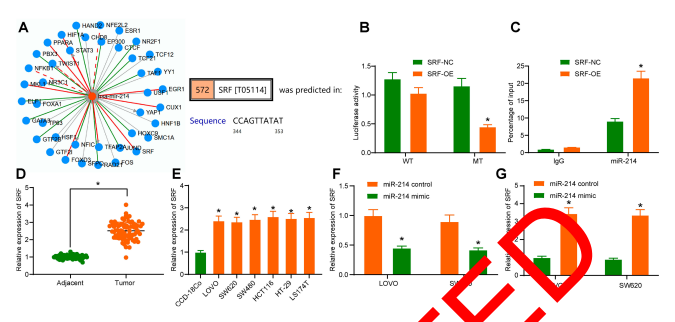


Figure 3 The binding relationship between miR-214 and SRF is identified. (A) the transcription factor the gulate miRand the ing promoter sequences predicted the effect of SRF-NC and SRF-OE on by TransmiR and ALGGEN; (B) the targeting relationship between SRF and miR-214 verified by a dual ase reporting a **A**PCR miR-214 promoter validated by ChIP; (D) SRF expression in CC and adjacent tissues evaluated by ction; (E) SRF pression in CC cell lines evaluated by RTqPCR detection; (F) SRF expression in CC cell lines overexpressing miR-214 evaluated by RT-qPCR detection; SRF expression in CC cell lines overexpressing miR-214 (panel BD. F evaluated by RT-qPCR detection. \*p < 0.05 according to one-way (panel (E) or two-way G). Data represent averages of three independent experiments.

## Knockdown of PTK6 Decreased CC Cell Viability

The viability detection of cells with PTK6 knock wn found that the number of EdU-positive cell as nota reduced (Figure 7A). Furthermore reduce PTK expression was linked to decreased ility tion (Figure 7B) and invasion (Figure 7C). importantly, CC cells with PTK6 knock n illustrate hampered tumorigenic and metastatic pote tials, supporting by a lighter tumor burder (figure 7D), Ner KI67 positive rate (Figure 7E) and smaller and fewer pulmonary nodules (Figure 7

# SRF/puR-21/PTK6 Axis Mediates the JAK2/STAT 3 Paulway

Finally, we explicitly the protein expression of JAK2 and STAT3 in cells cansfected with SRF-NC, SRF-OE, miR-214 control, miR-214 mimic, miR-214 mimic + SRF-NC, miR-214 mimic + SRF-OE, PTK6-NC or si-PTK6. Overexpression of SRF was found to expedite the JAK2/STAT3 signaling pathway, while miR-214 restoration and PTK6 knockdown diminished their phosphorylation. Besides, SRF flattened the inhibitory role of miR-214 mimic on the JAK2/STAT3 pathway induction (Figure 8A and B).

## iscussi

CC is as a result of genomic instability with pathering of genetic errors because of dysregulation of molecular signaling pathways governing cell migration, apoptosis and proliferation.<sup>23</sup> miRNAs have been indicated to play vital parts in the metastasis in CC. In the present investigation, miR-214 regulated by SRF was verified to inhibit metastasis and progression of CC by directly targeting the PTK6/JAK2/STAT3 pathway.

Previously, miR-214 was acknowledged as a tumor suppressor in the progression of cell migration, invasion and drug sensitivity of cervical cancer, implying its role as a promising diagnostic and therapeutic option for cancers.<sup>24</sup> The similar role of miR-214 has also been indicated in gastric cancer.<sup>25</sup> Moreover, circulating miR-214 has exhibited diagnostic potentials in breast cancer as an indicator of metastatic spread to lymph nodes.<sup>26</sup> As for its relevance in CC, miR-214 was notably diminished in the tissues from CC patients and cell lines, which was tightly linked to promoted cell proliferation.<sup>27</sup> Besides, miR-214 sensitized CC cells to irradiation by suppression of ATG12-regulated autophagy.<sup>28</sup> Even though miR-214 was corroborated to inhibit CC tumor growth in nude mice,<sup>29</sup> its connection with metastasis in vivo remains unclear. Our microarray analysis revealed that miR-214

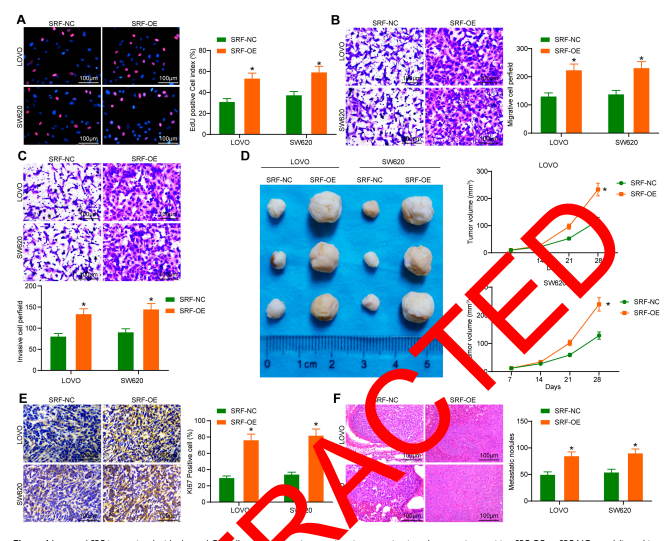


Figure 4 Increased SRF is associated with elevated C cell productive, migratory, invasive, tumorigenic and metastatic capacities. SRF-OE or SRF-NC was delivered into CC cell lines. (A) CC cell proliferation determined cell staining. CC cell migration evaluated by Transwell assays; (C) CC cell invasion evaluated by Transwell assays; (D) representative tumor images and tumor y min com mice injects with CC cells overexpressing SRF; (E) K167 positive rate of tumors detected by immunohistochemistry; (F) changes of pulmonary nodule detects by HE staining. p < 0.05 according to the two-way ANOVA. Data represent averages of three independent experiments.

was one of the most sufficant downregulated miRNAs niR-214 Apression profile in in CC. We further validate y La suggested that miR-CC tissue same es an cells. minished in both CC tissues and cell 214 expression was lines. Addited , this study exposed that miR-214 d the proliferative, migratory and invarestoration hamp sive capacities of Coccells, and most significantly, metastasis and tumor growth in vivo. Even though the biological functions of miR-214 knockdown have not been investigated in the current work, both Chandrasekaran et al and Zhou et al demonstrated that suppression of miR-214 increased proliferation, migration and invasion in CC cells and metastasis in vivo.<sup>30,31</sup>

To disclose the molecular mechanism of miR-214 in CC, we explored both the upstream and downstream

genes of miR-214. The bioinformatic tools revealed that the transcription factor SRF could bind to miR-214, and their binding relationship was substantiated by the following dual-luciferase and ChIP assays. SRF has been proposed to regulate the hepatic stellate cells.<sup>32,33</sup> Moreover, SRF and myocyte enhancer factor-2 worked collaboratively to modulate the expression of muscle-specific genes, such as miR-133a.<sup>34</sup> The myocar-din-related transcription factors A and B-SRF pathway is of great significance for cell proliferation, motility, and adhesion, which were all major processes underlying the progression of cancers.<sup>35</sup> For instance, SRF expedited gastric cancer metastasis by accelerating myo-fibroblast-cancer cell crosstalk.<sup>36</sup> More specifically, SRF was monitored to transactivate miR-199a-5p and miR-

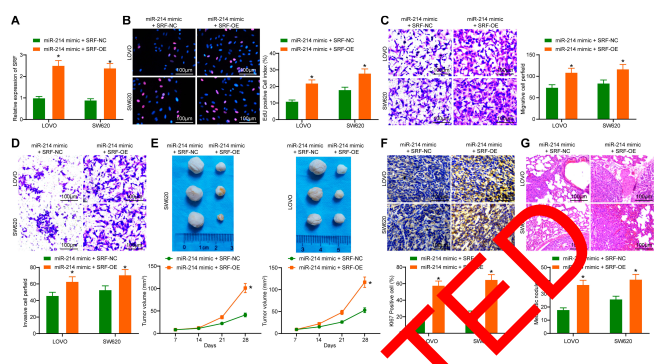


Figure 5 SRF overexpression rescues the repressive role of miR-214 mimic on CC cell viability CC cell lines were considered with SRF-OE or SRF-NC in the presence of miR-214 mimic. (A) SRF expression in CC cell lines co-transfected with miR-214 mimic at entropy evaluated by Rice SR detection; (B) CC cell proliferation determined by EdU staining; (C) CC cell migration evaluated by Transwell assays; (D) CC cell invasion evaluated by Transwell assays; (E) representative tumor images and tumor volume from mice injected with CC cells overexpressing SRF; (F), K167 positive rate of tumors extected by immunic histochemistry; (G) changes of pulmonary nodules detected by HE staining. \*p < 0.05 according to the two-way ANOVA. Data represent averages of the eindependent operiments.

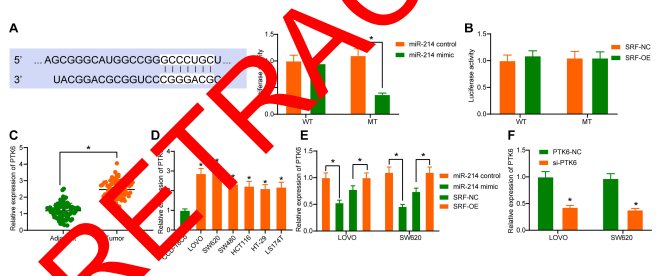


Figure 6 The target relationship between miR-214 and PTK6 is identified. (A) the targeting relationship between PTK6 and miR-214 predicted by bioinformatic analysis and verified by a relationship between PTK6 and SRF verified by a dual-luciferase reporting assay; (C) PTK6 expression in CC an ediacent tissues evaluated by RT-qPCR detection; (D) PTK6 expression in CC cell lines evaluated by RT-qPCR detection; (E) PTK6 expression in CC cell lines overexpressing miR-214 or SRF evaluated by RT-qPCR detection; (F) PTK6 expression in CC cell lines with PTK6 knockdown evaluated by RT-qPCR detection. \*p < 0.05 according to one-way (panel (D) or two-way ANOVA (panel AC, E and F). Data represent averages of three independent experiments.

199a-3p by interacting with their promoters in a direct manner to modulate epithelial-mesenchymal-transition and pulmonary metastases.<sup>37</sup> Similarly, we found that SRF overexpression enhanced CC cell metastatic and tumorigenic potentials by impeding the transcription of miR-214.

A miRNA prediction database revealed PTK6 as a candidate target of miR-214; we next carried out a dual-luciferase reporter assay and RT-qPCR to verify this hypothesis. Consistently, Cagle et al validated that miR-214 overexpression induced prostate cancer cell apoptosis, while repressed cell proliferation and colony

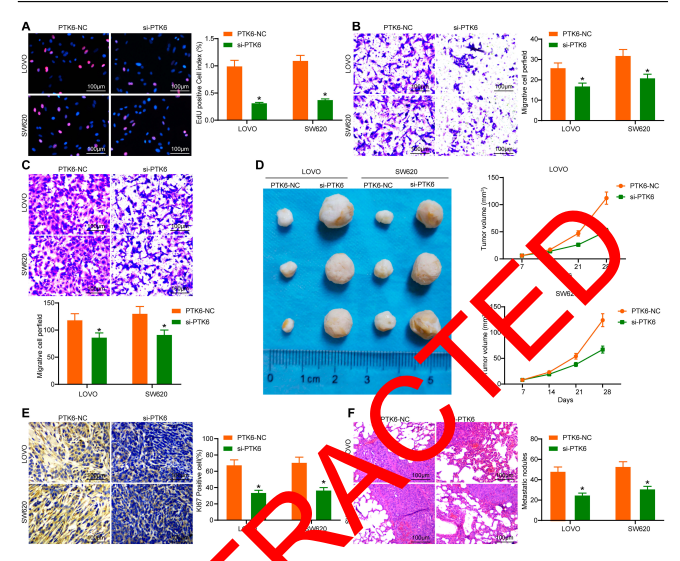


Figure 7 PTK6 knockdown is associated with rest of CC cell productive, migratory, invasive, tumorigenic and metastatic capacities. Si-PTK6 or PTK6-NC was delivered into CC cell lines. (A) CC cell proliferation domined by EdU stainine B) CC cell migration evaluated by Transwell assays; (C) CC cell invasion evaluated by Transwell assays; (D) representative tumor images and tumor value from mice injected with CC cells with PTK6 knockdown; (E) Kl67 positive rate of tumors detected by immunohistochemistry; (F) changes of tumonary nodule detected by HE staining. \*p < 0.05 according to the two-way ANOVA. Data represent averages of three independent experiments.

forming capacitie by theeting PTK6.38 PTK6 overngly prrelative the grade, recurrence expression st mosis of bladder cancer patients, as well as poor pr and PTK6 km arown contributed to considerably suppressed cell protoration and migration.<sup>39</sup> Meanwhile, the oncogenic role of PTK6 was also revealed by Wozniak et al in prostate cancer.<sup>40</sup> Furthermore, sustained induction of the JAK2/STAT3 pathway is highly correlated to tumorigenesis, proliferation, apoptosis and metastasis of tumors, including CC.<sup>41</sup> While PTK6 overexpression might induce the expression of STAT3 and proliferation of breast cancer cells.<sup>42</sup> In addition, miR-214 modulated CC procession through c-Met sialylation via the JAK2/STAT3 pathway.43 In the current

work, our Western blot results revealed that miR-214 overexpression or PTK6 knockdown reduced the extent of JAK2 and STAT3 phosphorylation, whereas SRF upregulation reversed the function of miR-214 on JAK2/STAT3 pathway activation.

All in all, the present study established that miR-214 downregulation, at least partially due to the upregulation of SRF, may promote CC growth and metastasis via activation of the PTK6/JAK2/STAT3 pathway, and miR-214 is probably a biomarker for predicting the prognosis of CC (Figure 9). Therefore, concentrating on the SRF/miR-214/PTK6/JAK2/STAT3 regulatory axis may be an attractive strategic option for the treatment of CC. Further examination is necessary for the validation of JAK2/STAT3 pathway in CC.

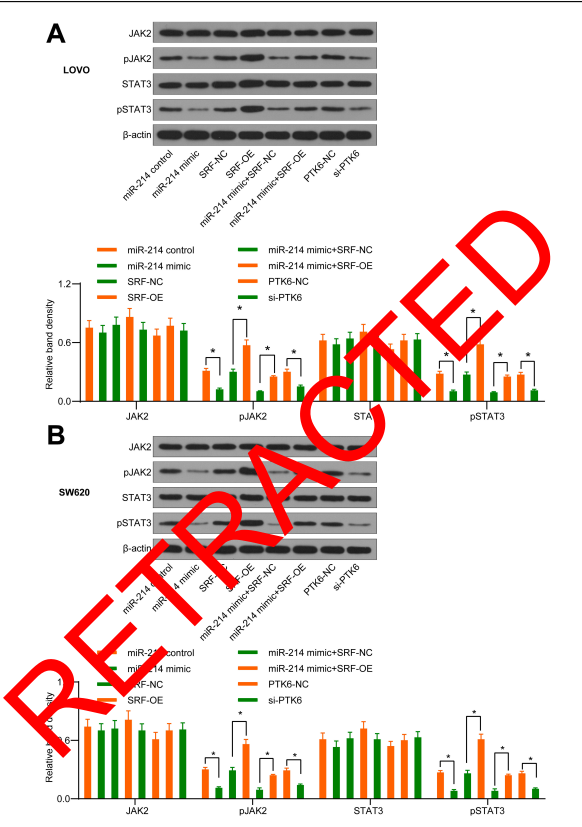


Figure 8 The JAK2/STAT3 pathway is regulated by the SRF/miR-214/PTK6 axis. The JAK2 and STAT3 protein expression and phosphorylation in LOVO (**A**) and SW620 (**B**) cells transfected with SRF-NC, SRF-OE, miR-214 control, miR-214 mimic, miR-214 mimic + SRF-NC, miR-214 mimic + SRF-OE, PTK6-NC or si-PTK6. \*p < 0.05 according to the two-way ANOVA. Data represent averages of three independent experiments.

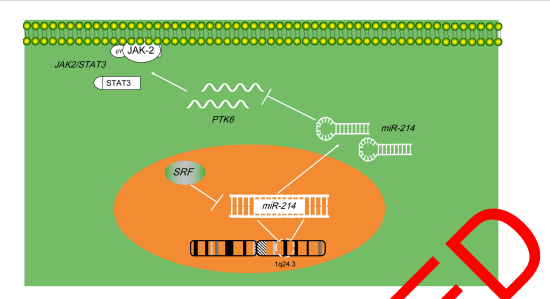


Figure 9 The model for miR-214 regulation upon CC cell progression. SRF could bind to the miR-214 promoter region, nated a chromosome 1 q24.3, and repress its expression. The restoration of miR-214 lowers the expression PTK6, thereby inhibiting the activity of the JAK2/ST 3 pathware mich in turn wibits CC cell proliferation, migration, invasion, tumorigenesis, and metastasis.

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## **Author Contributions**

TL contributed to the conception of the study V and ZYS designed the study. JYL provided adv e on s istical methods and the analyses of the data. NH the draft of the manuscript, tables d figu TL and draft of th CYZ provided comments on the manuscript. All authors contributed to data a lysis, drafting or revising the article, gave from approval of the version to be published, and agree to be accountable for all aspects of the work.

## Disclosure

The authors sla no con t of interest.

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