**Original Article**

**miR-601 inhibits proliferation, migration and invasion of prostate cancer stem cells by targeting KRT5 to inactivate the Wnt signaling pathway**

Hongbing Du, Xinghuan Wang, Rui Dong, Dongliang Hu, Yaoyi Xiong

*Department of Urology, Zhongnan Hospital of Wuhan University, Wuhan 430071, Hubei, P.R. China*

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**Abstract:** Objective: This study aimed to verify the hypothesis that downregulation of miR-601 inhibits the proliferation, migration, and invasion of prostate cancer stem cells (PCSCs) by the Wnt signaling pathway through targeting keratin 5 (KRT5). Methods: Bioinformatic tools were applied to predict miRNAs and genes potentially associated with prostate cancer, then miR-601 and KRT5 were selected. Subsequently, PCSCs were investigated with respect to miR-601 overexpression or inhibition, KRT5 overexpression, or treatment with a Wnt pathway inhibitor. A series of experiments including western blotting, RT-qPCR, wound healing experiment, transwell assay, MTT assay, annexin V-FITC/PI flow cytometric analysis, nude mice assay and immunohistochemistry were then carried out. Results: Compared with negative control group, migration, invasion, and proliferation of PCSCs and Wnt-1 expression were all enhanced, but apoptosis was attenuated in the miR-601 mimic group. Furthermore, results identified in the other groups (KRT5, miR-601 inhibitor, miR-601 inhibitor + KRT5, Wnt signaling pathway inhibitor, PRI-724/PRI-724 + KRT5) were opposite to those identified with the miR-601 mimic group (all \( P < 0.05 \)). Compared with the miR-601 inhibitor + KRT5 group, migration, invasion, and proliferation of PCSCs and Wnt-1 expression were all increased, whereas apoptosis was suppressed in KRT5 or miR-601 inhibitor groups (all \( P < 0.05 \)). Compared with the PRI-724 + KRT5 group, migration, invasion, and proliferation of PCSCs and Wnt-1 expression were also enhanced, whereas apoptosis was inhibited in PRI-724 or KRT5 groups (all \( P < 0.05 \)). Conclusion: Results obtained from the present study have demonstrated that downregulation of miR-601 is able to inhibit the proliferation, migration, and invasion of PCSCs by activating KRT5, and subsequently inhibiting the Wnt pathway.

**Keywords:** MicroRNA-601, keratin 5, prostate cancer, proliferation, migration, invasion, Wnt signaling pathway

**Introduction**

Prostate cancer is one of the most commonly diagnosed cancers in men, which affects the male reproductive system [1, 2]. Prostate cancer stem cells (PCSCs), which have the ability to self-renew and proliferate infinitely, have been reported to be the main source of prostate cancer [3]. To date, treatments for prostate cancer include resection, radiation therapy, and endocrine therapy [4, 5]. However, at present, no effective method is available for treating prostate cancer at an advanced stage. Gene therapy is considered to be one of the most promising treatments that may be utilized in efforts to cure prostate cancer, and this has become the most important component of composite treatments [6, 7]. The identification of novel therapeutic targets for more effective treatment of metastatic prostate cancer is urgently needed in order to improve disease management and patient survival.

MicroRNAs (miRNAs; miRs) have been reported to serve regulatory roles in the biologic and pathologic processes of various tumor types, including carcinogenesis, tumor growth, tumor differentiation, and cancer apoptosis [8, 9]. A previous study has demonstrated that miRNAs act either as tumor suppressors or as promoters by base-pairing with target genes and altering various signaling pathways associated with the disease under investigation [10]. Previous studies have indicated that miR-601 is involved in pancreatic cancer and colorectal cancer [11-13]. However, few studies to date have con-
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confirmed the function of miR-601 in prostate cancer. Keratin 5 (KRT5) is an intermediate filament protein expressed in the basal layer of stratified epithelial cells [14]. Accumulating evidence has indicated that KRT5 expression is downregulated in bladder cancer, and in head-and-neck squamous cell carcinoma [15, 16]. Therefore, the present study aimed to verify whether KRT5 is differentially expressed in prostate cancer by using a bioinformatics method, and it was subsequently confirmed that KRT5 is the target gene of miR-601. Wnt signaling pathway is an important molecular pathway that has a vital role in regulating the function of stem cells, and is reported to be an underlying cause of malignant cancers [17]. Therefore, a further aim of the present study was to explore whether miR-601 is able to affect the proliferation, migration, and invasion of PCSCs by regulating KRT5 and the Wnt signaling pathway. Taken together, the results of the present study have revealed that downregulation of miR-601 is able to inhibit the proliferation, migration, and invasion of PCSCs by activating KRT5, and subsequently inhibiting the Wnt pathway.

Materials and methods

Ethic statements

The use of tissues and all experimental procedures were carried out in accordance with the Declaration of Helsinki and approved by the ethics committee of Zhongnan Hospital of Wuhan University. All patients enrolled in the study signed informed consent documentation. All animal experiments took place at Zhongnan Hospital of Wuhan University, were approved by the local animal ethics committee, and followed the instructions of the Guide for Care and Use of Laboratory Animals published by the National Institutes of Health [18].

Bioinformatics prediction

A gene expression chip (GSE46602) of prostate cancer submitted by Aarhus University Hospital was downloaded from the Gene Expression Omnibus (GEO) database (http://www.ncbi.nlm.nih.gov/geo). Gene expression in prostate cancer tissue and the adjacent tissue of 36 cases of patients with prostate cancer was subsequently analyzed. Data normalization and background correction were performed using the Affy package (https://www.bioconductor.org/packages/devel/bioc/html/affy.html) based on the robust multiarray average algorithm [18]. Differentially expressed genes of prostate cancer were identified with R language of the limma package (https://master.bioconductor.org/packages/release/bioc/html/limma.html). Adjusted P-values were presented as adj.P.Val, and |logFC|>2.0 and adj.P.Val<0.01 were considered to represent the cutoffs for differentially expressed genes. The DisGeNET database (https://www.disgenet.org/web/DisGeNET/menu/search?4) was used to identify genes associated with prostate cancer. The potential interactions of the relevant genes were further mapped out using the STRING database (https://string-db.org) [19]. Subsequently, Cytoscape software (version 3.6.0) was adopted to visualize the differentially expressed genes and the disease interaction network. RNA22 (https://cm.jefferson.edu/ma22/), miPATH (http://lgmb.fmrp.usp.br/mirnopath/tools.php) and TargetScanHuman 7.2 (http://www.targetscan.org/vert_72/) were used as tools to predict the putative miRNA-mRNA interactions and the regulation of miRNAs with respect to their target genes. Venny 2.1.0 (http://bioinfogp.cnb.csic.es/tools/venny/index.html) was used to compare the differential genes identified in the chip assay and to construct Venn diagrams [20].

Study subjects

A total of 174 tissue samples (87 for prostate cancer tissue, and 87 for the adjacent tissues, respectively) of patients diagnosed at Zhongnan Hospital of Wuhan University between November 2015 and 2018 were involved in the present study. The mean age of the patients was 68.8 ± 7.75 (range: 36-69 years old). According to the tumor-node-metastasis (TNM) standard published by the American Joint Committee on Cancer (AJCC), 20 cases were confirmed to be at stage I, whereas 67 cases were at stage II. In terms of the degree of differentiation, 22 cases were classed as moderately or well differentiated, whereas 65 cases were poorly differentiated. Fifty-seven patients had lymph node metastasis, while 30 cases were free of any lymph node metastasis. The inclusion criteria were as follows [21]: i) the primary prostate cancer was clinically diagnosed and confirmed by pathologic examination; ii) the patient received no chemotherapy or radiotherapy prior to surgery; iii) the patient had indi-
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cations for surgical excision and could be treated by resection; iv) the survival time of the patient was >1 year. The exclusion criteria were as follows [22]: i) patients with acute prostatitis, benign prostatic hyperplasia, prostate cancer, neurogenic bladder, urethral deformity or stricture were excluded, as were those with: ii) heart, brain, liver or hematopoietic system disease; iii) mental illness, causing patients not to cooperate with the treatment; and iv) patients with severe complications during or after the operation.

Cell culture

The human prostate cancer cell line, PC-3, purchased from the ATCC (Manassas, VA, USA), was cultured with serum supplemented with RPMI-1640 medium with 5% CO₂ at 37°C. When the cell confluence had reached 80%, the cells were treated with 0.25% trypsin, and cells in the exponential growth phase were selected for subsequent experiments.

Preparation and confirmation of PCSCs

The PC-3 cells in the exponential growth phase were resuspended in serumfree medium (SFM), and the cell concentration was adjusted to 1×10³ cells/ml. Subsequently, the cells were seeded in a 100 cm² plate at a density of 1×10³ cells per plate, and the plate was agitated every day during the incubation to avoid cell attachment. After the cells had grown and formed spheres, they were removed and placed in a centrifuge tube, and almost half the supernatant was discarded 20 min after their placement, followed by addition of an equivalent amount of SFM to separate the cells and form a single cell suspension. The cells were subsequently resuspended in SFM and subcultured at a ratio of 1:2 in an incubator with 5% CO₂ at 37°C. An inverted microscope (Olympus CKX31; Olympus Corporation, Tokyo, Japan) was used to observe the cells at 5, 10 and 15 days after seeding. Flow cytometry analysis was subsequently utilized to detect the proportion of the CD133⁺CD44⁺ cells in the PC-3 cell spheres and in adherent cells. These two types of cells were resuspended to form a single cell suspension; 2 ml cell suspension (with a concentration of 1×10⁴ cells/ml) was then collected and stained using anti-CD44-FITC (cat. no. ab27285; Abcam, Cambridge, UK) and anti-CD133-PE (cat no. 130-098-826; Beijing Beads Biotechnology Co., Ltd., Beijing, China) antibodies. After incubation on ice in the dark for 30 min and washing with PBS for 3 times, flow cytometry (using a CyFlow Cube6 flow cytometer; Sysmex Partec, Görlitz, Germany) was adopted to measure the apoptotic rate of the CD44⁺CD133⁺ cells.

Construction of the KRT5 expression vector

The primer sequence of KRT5 mRNA (accession number: NM_000424.3) was obtained from the nucleotide database of NCBI, and subsequently synthesized by Shanghai Gencem Co., Ltd. (Shanghai, China). The correctly synthesized sequence was cloned into Invitrogen™ plasmid vector pcDNA3.1 (+) (VPI0001; Thermo Fisher Scientific, Inc., Waltham, MA, USA) after HindIII and XhoI digestion and ligation had been performed with T4 ligase at 16°C for 1 h. Subsequently, the ligation products were transferred to DH5α competent cells (d9052; Takara Biotechnology Ltd., Dalian, China), and resistant colonies of the bacteria were selected and identified by reverse transcriptionquantitative polymerase chain reaction (RT-qPCR), as described below. Finally, the plasmids were extracted using a PicoPure® DNA Extraction kit (KIT0103; Thermo Fisher Scientific, Inc.) and stored at -20°C before use.

Dual-luciferase reporter gene assay

MiR-601 target genes were further predicted using the online miRNA prediction tool, Targetscan (http://www.targetscan.org/vert_72/). KRT5 was identified as a potential miR-601 direct target gene, and subsequently a dual-luciferase reporter assay was utilized for experimental validation. The firefly luciferase reporter plasmid containing a KRT5-3'-UTR-wt (or KRT5-3'-UTR-mut) sequence was established, and these were respectively cotransfected along with miR-601 (miR-601 NC) into CHO cells (CC-Y2110; ATCC, USA). Renilla luciferase plasmid served as a control. Fluorescence intensity was measured using a dual-luciferase reporter gene assay kit (GM-040502A; Products, China) under a DFM-20 fluorescence microscope (Cai Kang Optics, China) at 560 nm (for firefly luciferase) or 465 nm (for Renilla luciferase), and the ratio of the firefly luciferase activity value/Renilla luciferase activity value was used to calculate the relative luciferase activity. Luciferase mRNA expression levels were determined using
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RT-qPCR, as described below. Experiments were repeated 3 times.

Cell grouping and transfection

The PCSCs were divided into the following groups: The negative control group (transfected with empty vector), miR-601 mimic group (transfected with miR-601 mimic), miR-601 inhibitor group (transfected with miR-601 inhibitor), KRT5 group (transfected with KRT5 overexpression recombinant plasmids), miR-601 mimic + KRT5 group (transfected with miR-601 mimic and KRT5 overexpression recombinant plasmids), PRI-724 group (treated with the Wnt signaling pathway inhibitor, PRI-724) and the PRI-724 + KRT5 group (treated with PRI-724 and transfected with KRT5 overexpression recombinant plasmids).

Cell transfection was performed as follows: Specifically, cells were seeded into a 20-well plate. Cell transfection was then conducted using an Invitrogen Lipofectamine 2000 kit (Thermo Fisher Scientific, Inc), following the manufacturer’s protocol. Aliquots (20 pmol) of KRT5 overexpression recombinant plasmids, miR-601 mimic, miR-601 inhibitor or scrambled control were dissolved in 50 µl PBS (designated as solution A), or they were further mixed with 50 µl PBS containing 1 µl Lipofectamine™ 2000 (solution B), and the solutions were placed at room temperature for 20 min. Subsequently, solutions A and B were mixed and added into cells, which were then cultured in an incubator with 5% CO₂ at 37°C. The medium was completely changed after incubation for 6-8 h.

The PRI-724 inhibitor (S8262; Selleck Chemicals, Houston, TX, USA) was diluted with serum-free medium SFM to 150 nM; subsequently, 2 ml of the mixture was added to the cells and incubated for a 24 h period, after which the culture medium was subsequently changed.

RT-qPCR

Total RNA was extracted from PCSCs featured in all the experimental treatment groups using a miRNeasy Mini kit (Tiangen Biotech Co., Ltd, Beijing, China) after 48 h transfection. RNA samples (5 µl) were diluted with 20× free-RNA enzyme ultrapure water. The optical density (OD) values (at 260 and 280 nm) and RNA concentration were detected using ultraviolet spectroscopy with a UV1901 double beam spectrophotometer (Aoxi Scientific Instrument Ltd., Shanghai, China). A ratio of OD_{260}/OD_{280} ranging from 1.7-2.1 was considered to indicate high purity, and these RNA samples were therefore selected for subsequent experiments. The reverse transcription process was performed using EasyScript® First-Strand cDNA Synthesis SuperMix (AE301-02; TransGen Biotech Company, Beijing, China), according to the manufacturer’s protocol. The thermocycling conditions of this procedure were as follows: 37°C for 15 min, followed by 85°C for 5 sec; the samples were then frozen at -80°C for later use. The primers of miR-601, KRT5, Wnt-1, β-catenin, Nanog, and Oct-4 were designed and synthesized by Sangon Biotech (Shanghai) Co., Ltd. (Shanghai, China) (Table 1).

The qPCR reaction was performed using a SYBR® Premix Ex Taq™ II kit (RR820A; Xing Zhi Biotech Co., Ltd., Guangzhou, Guangdong Province, China), according to the manufacturer’s protocol. The thermocycling conditions of this procedure were as follows: 37°C for 15 min, followed by 85°C for 5 sec; the samples were then frozen at -80°C for later use. The primers of miR-601, KRT5, Wnt-1, β-catenin, Nanog, and Oct-4 were designed and synthesized by Sangon Biotech (Shanghai) Co., Ltd. (Shanghai, China) (Table 1). The qPCR reaction was performed using a SYBR® Premix Ex Taq™ II kit (RR820A; Xing Zhi Biotech Co., Ltd., Guangzhou, Guangdong Province, China), according to the manufacturer’s protocol. The reaction system included 10 µl SYBR Premix, 2 µl cDNA template, 0.6 µl forward and reverse primers, and 6.8 µl sterile water. RT-qPCR was performed with an ABI 7500 Real-Time PCR system (ABI Research, Oyster Bay, NY, USA), and glyceraldehyde phosphate dehydrogenase (GAPDH) and U6 were used as internal references for the genes of interest and miR-601, respectively. The thermocycling conditions were as follows: pre-denaturation at 95°C for 30 sec, 45 cycles of

| Table 1. Primer sequences for quantitative real-time polymerase chain reaction |
|------------------|------------------|
| Gene             | Primer Sequences |
| KRT5             | 5’-CCAAGGGTGGTACGCTGATGG-3’<br>5’-GTGACGACATGTCGCTG-3’ |
| Wnt1             | 5’-TTCAGACAGAGATGGAAGACT-3’<br>5’-CCACGGCTCAGTGGAG-3’ |
| Oct4             | 5’-GGAGAGAACATGGAAGCT-3’<br>5’-AAAGCTGAGAAATCCCTG-3’ |
| β-catenin        | 5’-ATGGAACAGAGAAGAC-3’<br>5’-AGGACTGAGAAATCCCTG-3’ |
| Nanog            | 5’-CTTCGGAGAGAAAGATTAGTCAG-3’<br>5’-AGTGGGACCTGTTAATCT-3’ |
| miR-601          | 5’-CAGTATGATTTGACTCTTTG-3’<br>5’-GTGACGGAAGCTGAGAGCT-3’ |
| U6               | 5’-AAAGCGATATCCGACGGACC-3’<br>5’-GTACACACATTTTCTCTGGA-3’ |
| GAPDH            | 5’-TGTGTCAGATGATGATGAG-3’<br>5’-ACACCATGATTCGCGGTCAAT-3’ |
denaturation at 95°C for 5 sec, annealing at 20°C for 30 sec and extension at 72°C for 30 sec. The $2^{ΔΔCq}$ method was utilized to calculate the relative mRNA expression levels of miR-601, KRT5, Wnt-1, β-catenin, Nanog and Oct4, according to the equation: $ΔΔCq = ΔCq_{\text{experimental group}} - ΔCq_{\text{control group}}$ [23]. The experiment was repeated 3 times.

**Western blot analysis**

PCSCs in the exponential growth phase were selected after 48 h and washed 3 times with pre-cooled PBS. Subsequently, the PCSCs were lysed on ice using RIPA buffer (R0010; Beijing Solarbio Science and Technology Co., Ltd., Beijing, China) for 30 min, centrifuged at 12,000 rpm for 10 min to obtain the supernatant, and stored at -20°C prior to determination of the protein concentration using a BCA kit (PC0020; Beijing Solarbio Science and Technology Co., Ltd.). The protein was subsequently treated with the same volume of SDS-PAGE sample buffer (P1200; Beijing Solarbio Science and Technology Co., Ltd.) in accordance with the manufacturer’s protocol. Subsequently, the proteins were transferred to a polyvinylidene fluoride membrane and blocked for 1 h with 5% BSA at room temperature. The membranes were then incubated overnight at 4°C after incubating with the primary antibodies against KRT5 (cat. no. ab52635, 1:10,000 dilution), Wnt-1 (cat. no. ab15251, 1:100), Oct4 (cat. no. ab18976, 1 μg/ml), β-catenin (cat. no. ab15180, 1:100), phosphorylated (p)-β-catenin (cat. no. ab27798, 1:500), Nanog (cat. no. ab106465, 1 μg/ml), matrix metalloproteinase 2 (MMP2) (cat. no. ab37150, 1 μg/ml), the tumor metastasis suppressor KAI1 (cat. no. ab59509, 1 μg/ml), N-cadherin (cat. no. ab18203, 1 μg/ml), E-cadherin (cat. no. ab-15148, 1:500), Bax (cat. no. ab32503, 1:1,000), Bcl2 (cat. no. ab32124, 1:1,000), and GAPDH (cat. no. ab37168, 1 μg/ml) (all purchased from Abcam). The next day, the membranes were washed with PBST buffer 3 times (10 min each wash), and subsequently incubated with the secondary antibody, goat anti-rabbit IgG (cat. no. ab205718, 1:5,000, Abcam), gently agitated for 1 h on a shaking table, and washed 3 times again with PBST (15 min each wash). The membranes were then immersed in electrochemiluminescence (ECL) solution (Pierce; now a brand of Thermo Fisher Scientific, Inc.) for 1 min, prior to exposure to X-rays in a dark room to develop and determine the color. Finally, the images were developed using a Bio-Rad gel imaging system (MG8600; Beijing Thmorgan Biological Technology Co., Ltd, Beijing, China), and the relative protein level was evaluated using ImageJ software. Each experiment was repeated 3 times.

**Wound healing experiment**

PCSCs were seeded into a 24-well plate (4×10^5 cells each well). After 24 h, a scratch was generated in each well by using a 200 μl sterile gun. The cells were washed 3 times with PBS, and then cultured in an incubator. The migration distance was detected using an inverted microscope (CX23; Olympus Corporation), and images were captured at 0 and 48 h after the scratch test. The healing rate was calculated according to the following formula: [migration distance (0 h) - migration distance (48 h)] / migration distance (0 h) [24]. Each experiment was repeated 3 times.

**Transwell assay**

Matrigel™ (BD Biosciences, San Jose, CA, USA) was diluted with precooled and serum-free DMEM medium in a ratio of 1:10. Subsequently, 100 μl diluted Matrigel was added to the upper chamber and placed at room temperature for 2 h. PCSCs in each group were washed with 200 μl SFM (comprising RPMI-1640 medium) and digested with 0.25% trypsin, before being resuspended with SFM (comprising DMEM) 24 h after transfection. Subsequently, 100 μl cells (adjusted to 3×10^5 cells/ml) were added into the Transwell upper chamber (Corning Inc., Corning, NY, USA), and 600 μl 10% serum comprising DMEM medium was added into the lower chamber at the same time. The experimental procedure was performed according to the manufacturer’s protocol. Three visual fields were randomly picked, and the number of cells crossing the membrane was counted under an inverted microscope (XDS-800D; Shanghai Caikon Optical Instrument Co., Ltd., Shanghai, China). The invasion rate was calculated according to the following equation: Invasion rate = (number of cells passing through the matrix/total cells) × 100% [25]. The experiment was repeated 3 times.

**Methyl thiazolyl tetrazolium (MTT) assay**

Single cell suspension was prepared by adding 10% fetal bovine serum to the cells, which
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were subsequently added to a 96-well plate [cell density, (3-6) ×10^3 cells per well, in a volume of 200 μl per well]. Six wells in duplicate were set up, and to each well was added 10 μl MTT solution (M1025; Beijing Solarbio Science and Technology Co., Ltd.) at 0, 24, 48 and 72 h after transfection. The cells were then cultured for a further 4 h. Dimethyl sulfoxide (DMSO) (150 μl) was then added each well, and the plate was agitated for 10 min to fully dissolve the crystals. An MTT enzyme-linked immunometric meter (DNM-9602G; Beijing Pulang Instrument Co., Ltd, Beijing, China) was adopted for OD detection at 490 nm. Each experiment was repeated 3 times.

**Annexin V-fluorescein isothiocyanate (FITC)/PI double staining**

An annexin V-FITC/PI double staining kit (556547; Becton, Dickinson and Company, USA) was used to measure apoptosis of the PCSCs following transfection for 48 h. Specifically, cells were treated with 0.25% trypsin, the cell density was adjusted to 1×10^6 cells/ml, and the cells in each group were subsequently centrifuged at 4,000 rpm/min for 5 min. Subsequently, the cells were collected and resuspended in 1×PBS, centrifuged again at 2,000 rpm for 5 min, and 300 μl 1× Binding Buffer was then added to the cells. Subsequently, 5 μl Annexin V-FITC was added to the resuspended cells, which were then incubated at room temperature for 15 min prior to the addition of 5 μl PI. A CyFlow Cube6 flow cytometer (Sysmex Partec) was used to determine the extent of apoptosis: The absorbances of FITC (at 480 nm and 530 nm) and PI (at 575 nm) were detected, respectively, at the different excitation wavelengths. This experiment was repeated 3 times.

**Tumor xenograft in nude mice**

In total, 35 BALB/C nude mice (weighing 18-25 g, and aged 4 weeks) were purchased from Hubei Research Center of Laboratory Animal (Wuhan, China) and raised in a specific pathogen environment. The mice were divided into seven groups (n = 5 mice in each group, which were set up to be the same as the cell group treatment groups. Subsequently, the mice were administered 2% sodium pentobarbital (0.6 ml/100 g) by intraperitoneal injection, and 0.2 ml PCSC suspension (concentration of cells, 10^7/ml) was inoculated into the subcutaneous tissue. At the end of the fourth week, the mice were sacrificed, and the resultant tumors were dissected and weighed. The tumor volume was calculated according to the formula: Length × width^2/2 [26].

**Immunohistochemical staining**

Lymph nodes of nude rats were obtained, and immediately fixed with 4% neutral buffered formalin (Beijing Solarbio Science and Technology Co., Ltd.) for 24 h. After washing with PBS, the lymph node tissues were dehydrated, cleared, embedded in paraffin, and sliced (3 μm slices). Before performing the experiments, these slices were baked at 60°C for 30 min, dewaxed in xylene (I and II), dehydrated through an ethanol gradient (100, 85 and 80%) for 5 min at each concentration, and washed with PBS for 3 times. Subsequently, antigen retrieval was accomplished through microwave treatment in citrate buffer (pH = 8.0) for 10 min, and the tissue slices were then washed 3 times again with PBS (5 min each wash) after cooling to room temperature. Subsequently, 0.3% H_2O_2-methanol solution was added to the slices at room temperature for 10 min to block endogenous peroxidase activity, prior to washing twice with PBS (5 min each wash). Subsequently, the slices were incubated with Ki-67 primary antibody (diluted to 1 μg/ml, cat. no. ab15580, Abcam) at 4°C overnight in a fridge, and washed three times with 0.1% PBST (5 min each wash) the following day. Polymer reinforcer was then added to the slices, which were incubated at room temperature for 20 min and washed three times again with PBST (5 min each wash). Horseradish peroxidase-labeled IgG, as a secondary antibody (PV-9000-D; Beijing Zhongshan Golden Bridge Biotechnology Co. Ltd., Beijing, China), was then added. The slices were subsequently incubated at room temperature for 30 min, and washed three times with 0.1% PBST (5 min each wash). Finally, the slices were stained with 3,3'-diaminobenzidine (DAB) for 5 min, re-stained with hematoxylin, and observed under an inverted microscope. IPP7.0 software (Media Cybernetics, Singapore) was used to perform semi-quantitative analysis. The cells were counted in 10 fields (100 PCSCs in each field) and the percentages of positively-stained cells were calculated using the formula: No. of positively-stained cells/total number of cells ×100% [27].
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Statistical analysis

Statistical analysis was performed using SPSS v. 21.0 software (IBM Corp., Armonk, NY, USA). Results were expressed as the mean ± standard deviation (SD). All experiments were repeated 3 times. Comparisons between two groups were analyzed using the t-test, whereas multiple groups were analyzed using one-way analysis of variance (ANOVA). The test of normality was verified by the Kolmogorov-Smirnov test. Specifically, the normal distribution in multiple groups was conducted by one-way ANOVA followed by the Tukey post-hoc test, whereas non-normal distribution analysis was analyzed using Dunn’s multiple comparison for post-hoc tests following Kruskal-Wallis testing. P<0.05 was considered significant.

Results

Bioinformatics prediction

Based on the information obtained from the prostate cancer gene chip, GSE46602, 153 differentially expressed genes were selected according to the set criteria, |logFC|>2.0 and adj.P.Val<0.01. The top 10 genes (KLK3, AR, ERG, TP53, PTEN, PROS1, NPEPPS, PSAT1, PLAG1, and TMPRSS2) reported to be associated with prostate cancer (code: C0600139) were screened from the database, DisGeNET (Figure 1A). Both the 10 differentially expressed genes and the target miRNA through using a bioinformatics analysis. A. Differentially expressed genes indicated in the chip data (GSE46602) associated with prostate cancer. The right panel is color-coded, where each rectangle represents a sample. The red color indicates high expression, whereas the blue color indicates relatively low expression. B. Gene interaction network of prostate cancer-associated genes and known genes. The red rectangles represent known disease genes, whereas the green rectangles represent differentially expressed genes. C. Comparison of RNA22, miRNApath and Targetscan databases on KRT5 miRNA upstream identified hsa-miR-601 and hsa-miR-564 as the intersecting miRNAs. KRT5, keratin 5.

Figure 1. Screening of prostate cancer-associated genes and the target miRNA through using a bioinformatics analysis. A. Differentially expressed genes indicated in the chip data (GSE46602) associated with prostate cancer. The right panel is color-coded, where each rectangle represents a sample. The red color indicates high expression, whereas the blue color indicates relatively low expression. B. Gene interaction network of prostate cancer-associated genes and known genes. The red rectangles represent known disease genes, whereas the green rectangles represent differentially expressed genes. C. Comparison of RNA22, miRNApath and Targetscan databases on KRT5 miRNA upstream identified hsa-miR-601 and hsa-miR-564 as the intersecting miRNAs. KRT5, keratin 5.
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miR-601 is predicted to target KRT5, and confirmation of this hypothesis by luciferase reporter gene assay

Through using the online prediction tool Targetscan, the binding sites of miR-601 and KRT5 were identified, and these are presented in Figure 3A. Dual luciferase report assay was utilized to further confirm that miR-455-3p directly mediates regulation of KRT5. The results also revealed that miR-601 could significantly reduce the luciferase activity of the KRT5-3′-UTR-wt, although it exerted little influence on KRT5-3′-UTRmut. These results confirmed that miR-601 directly targets KRT5 (Figure 3B) (P<0.05).
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Preparation of prostate cancer cell spheres

After culture in SFM for 4-5 days, the stem cell spheres were initially formed, and these demonstrated good refraction and high density; subsequently, both their size and density increased after 10-15 days. The results of the flow cytometric analysis revealed that the percentage of CD133^+ CD44^+ cells was 81.26% in the stem cell spheres, much higher compared with that in the monolayer attached cells (4.15%) (Figure 4). The stem cell spheres were selected for the subsequent cell experiments with PCSCs.

MiR-601 participates in activation of the WNT signaling pathway by targeting KRT5

As shown in Figure 5, the results of RT-qPCR and western blot analysis demonstrated that, compared with the negative control group, the miR-601 mimic group exhibited a reduced level of expression of p-β-catenin and KRT5, but an increased expression level of miR-601, Wnt-1 and β-catenin were observed, whereas the miR-601 inhibitor and KRT5 groups demonstrated the opposite results (all \( P < 0.05 \)). Compared with the miR-601 inhibitor + KRT5 group, the expression levels of p-β-catenin and KRT5 were reduced, whereas that of Wnt-1 and β-catenin were increased, in the KRT5 and miR-601 inhibitor groups (all \( P < 0.05 \)).

Inhibition of miR-601 or overexpression of KRT5 inhibits the migration and invasive abilities of PCSCs by regulating the Wnt signaling pathway

Western blotting was utilized to determine the expression levels of proteins associated with invasion and metastasis (Figure 6). These results indicated that, compared with the negative control group, the protein expression levels of N-cadherin and MMP-2 in the miR-601 mimic group were upregulated, but these proteins were downregulated in the other groups (all \( P < 0.05 \)). Furthermore, the protein expression levels of the tumor metastasis suppressor, KAI1, and E-cadherin in the miR-601 mimic group were clearly decreased, but were increased in the other groups (all \( P < 0.05 \)). Compared with the miR-601 inhibitor + KRT5 group, the protein expression levels of N-cadherin and MMP-2 in the miR-601 inhibitor and KRT5 groups were increased, whereas...
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The results of the wound healing and Transwell assays indicated that, compared with the PCSCs in the negative control group, the migration and invasive capabilities of the PCSCs in the miR-601 mimic group were significantly enhanced, although these were diminished in the other groups (all \(P<0.05\)). Compared with the miR-601 inhibitor + KRT5 group, the migration and invasive abilities of PCSCs in the KRT5 and miR-601 inhibitor groups were significantly strengthened (all \(P<0.05\)). It was also noted that the migration and invasive capabilities of the PCSCs in the KRT5 and PRI-724 groups were significantly reinforced in comparison with the PRI-724 + KRT5 group (all \(P<0.05\)) (Figures 7 and 8).

Inhibition of miR-601, or overexpression of KRT5 inhibits PCSC proliferation by the Wnt signaling pathway

Western blotting was employed to measure the levels of proteins associated with proliferation, and the results (Figure 9) revealed that, compared with the negative control group, the protein expression of Nanog and Oct4 was upregulated in the miR-601 mimic group, but downregulated in all other groups (all \(P<0.05\)). Compared with the miR-601 inhibitor + KRT5 group, the protein expression levels of Nanog and Oct4 in the miR-601 and KRT5 groups were increased (all \(P<0.05\)). In addition, compared with the PRI-724 + KRT5 group, the protein expression levels of Nanog and Oct4 in the PRI-724 and KRT5 groups was increased (all \(P<0.05\)).

Figure 5. Activation of the Wnt signaling pathway. A. RT-qPCR was utilized to detect the expression levels of miR-601 and mRNA as they relate to the Wnt signaling pathway. B. Protein expression levels of proteins associated with the Wnt signaling pathway were determined by western blot analysis. C. Quantification of the protein expression data is shown. \(^*P<0.05\) cf. the negative control group; \(^\#P<0.05\) cf. the miR-601 inhibitor + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5.

Figure 6. Relative protein levels of MMP-2, KAI1, N-cadherin and E-cadherin. A. Protein expression levels of MMP-2, KAI1, N-cadherin and E-cadherin were determined by western blot analysis. B. Quantification of the protein expression is shown. \(^*P<0.05\) cf. the negative control group; \(^\#P<0.05\) cf. the miR-601 inhibitor + KRT5 group; \(^&P<0.05\) cf. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5; MMP-2, matrix metalloproteinase 2.
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Figure 7. Migration ability of the PCSCs in each group as detected by scratch wound assay. A. Maps of the cell scratch test are shown. B. Quantification of the wound closure in each group is shown. *P<0.05 cf. the negative control group; †P<0.05 cf. the miR-601 inhibitor + KRT5 group; ‡P<0.05 cf. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5; PCSC, prostate cancer cell line.

Figure 8. Invasive ability of PCSCs in each group as determined by Transwell assay. A. Observation of the invasion of PCSCs in each group. B. Quantification of the invasiveness of PCSCs is shown. *P<0.05 cf. the negative control group; †P<0.05 cf. the miR-601 inhibitor + KRT5 group; ‡P<0.05 cf. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5; PCSC, prostate cancer cell line.

MTT assay was employed to detect the proliferative ability of the PCSCs in each group. As a result, no significant differences in cell proliferation were observed comparing among the groups during the 0-24 h time period (all P<0.05). Compared with the negative control group, however, cell proliferation was significantly enhanced following miR-601 overexpression, although the cell proliferative ability was reduced in the other groups (all P<0.05). Compared with the miR-601 inhibitor + KRT5 group, the proliferative ability of the PCSCs in the KRT5 and miR-601 inhibitor groups was significantly strengthened (all P<0.05). Finally, the proliferative ability in the KRT5 and PRI-724 groups was significantly increased in comparison with the PRI-724 + KRT5 group (all P<0.05) (Figure 10).

Inhibition of miR-601 or KTR5 overexpression inhibits apoptosis of PCSCs by the Wnt signaling pathway

Western blotting was performed to measure the levels of the apoptosis-associated proteins, Bcl-2 and Bax, in each group, and these results (Figure 11) showed that, compared with the
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Figure 9. Relative protein levels of Nanog and Oct4. A. Protein expression levels of Nanog and Oct4 were determined by western blot analysis. B. Quantification of the protein expression data is shown; *P<0.05 vs. the negative control group; **P<0.05 vs. the miR-601 inhibitor + KRT5 group; ***P<0.05 vs. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5.

Figure 10. Proliferative ability of the PCSCs in each group as determined by MTT assay. Proliferative ability of the PCSCs was investigated through using the MTT assay at 0, 24, 48 and 72 h. *P<0.05 vs. the negative control group; **P<0.05 vs. the miR-601 inhibitor + KRT5 group; ***P<0.05 vs. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5; PCSC, prostate cancer stem cell.

The cellular biological experiments described above were performed first, and the results obtained revealed that miR-601 was able to regulate the biological characteristics of PCSCs by regulation of the Wnt signaling pathway by targeting KRT5. Subsequently, xenograft tumor experiments in nude mice were performed to further explore the involvement of miR-601 in the regulation of PCSCs through interactions with KRT5 and the Wnt signaling pathway. These results demonstrated that, compared with the negative control group, the gross tumor volume in the miR-601 mimic group was significantly increased, although this was decreased in the other groups (all P<0.05). Compared with the miR-601 inhibitor + KRT5 group, the tumor volume in the KRT5 and miR-601 inhibitor groups was significantly suppressed (all P<0.05). Finally, the levels of apoptosis in the KRT5 and PRI-724 groups were significantly downregulated in comparison with the PRI-724 + KRT5 group (all P<0.05) (Figure 12).

Inhibition of miR-601, or overexpression of KTR5, inhibits tumor growth in nude mice by regulation of the Wnt signaling pathway

The cellular biological experiments described above were performed first, and the results obtained revealed that miR-601 was able to regulate the biological characteristics of PCSCs by regulation of the Wnt signaling pathway by targeting KRT5. Subsequently, xenograft tumor experiments in nude mice were performed to further explore the involvement of miR-601 in the regulation of PCSCs through interactions with KRT5 and the Wnt signaling pathway. These results demonstrated that, compared with the negative control group, the gross tumor volume in the miR-601 mimic group was significantly increased, although this was decreased in the other groups (all P<0.05). Compared with the miR-601 inhibitor + KRT5 group, the tumor volume in the KRT5 and miR-601 inhibitor groups was significantly suppressed (all P<0.05). Furthermore, the tumor volumes in the KRT5 and PRI-724 groups were clearly larger compared with the PRI-724 + KRT5 group (all P<0.05) (Figure 13).

Inhibition of miR-601 or KTR5 overexpression inhibits lymph node metastasis by regulating the Wnt signaling pathway

To detect lymph node metastasis in each group, immunohistochemistry was performed to measure the protein level of Ki-67. It was found
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Figure 11. Relative protein levels of Bax and Bcl-2. A. Protein expression levels of Bax and Bcl-2 were determined by western blot analysis. B. Quantification of the protein expression data is shown. *P<0.05 vs. the negative control group; #P<0.05 vs. the miR-601 inhibitor + KRT5 group; &P<0.05 vs. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5.

that, compared with the negative control group, the positive rate of Ki-67 protein expression in the miR-601 mimic group was significantly upregulated; whereas, in the other groups, the positive rate of Ki-67 was downregulated (all P<0.05). Compared with the miR-601 inhibitor + KRT5 group, the expression levels of Ki-67 protein in the KRT5 and miR-601 inhibitor groups were significantly increased (all P<0.05). The positive rate of Ki-67 protein in the KRT5 and PRI-724 groups was also identified to be markedly increased in comparison with the PRI-724 + KRT5 group (all P<0.05) (Figure 14).

Discussion

Prostate cancer is one of the most commonly diagnosed cancers in men [39]. Pharmacotherapy, immunotherapy, radiotherapy, chemotherapy, and palliative therapy are the common methods of treatment [40]. From the perspective of gene therapy, the present study aimed to explore the role of miR-601-KRT5 interaction on the Wnt signaling pathway in the pathogenesis of prostate cancer. As a result, it has been demonstrated that the inhibition of miR-601 may activate KRT5 and suppress the Wnt signaling pathway, and subsequently, positively affect the proliferation, migration, and invasion of PCSCs.

Initially, bioinformatic tools were employed to predict potential genes which are associated with prostate cancer, and it was revealed that the recurrence and chemotherapeutic resistance of ovarian cancer. All these findings suggested a positive role of KRT5 in cancer. miR-601 is the target RNA of KRT5, and a study reported that miR-601 overexpression could serve as an important marker in the diagnosis of colorectal cancer [34]. However, the role of miR-601 in prostatic cancer remains largely unknown. In order to explore the effects of miR-601 and KRT5 on PCSCs, real-time fluorescence quantitative assay and western blot assays were performed, and the results obtained demonstrated that the inhibition of miR-601, or KRT5 overexpression, are able to inhibit the expression of Nanog and Oct4, and that the miR-601 inhibitor + KRT5 group exhibited the most obvious decline. In addition, the expression of KRT5 was significantly decreased, whereas the expression levels of Wnt1 and β-catenin were significantly increased in the miR-601 mimic group, suggesting that the effects of miR-601 on PCSCs is mainly mediated through its interaction with KRT5, and by regulation of the Wnt signaling pathway. The Wnt signaling pathway exerts a crucially important role in the pathogenesis of prostate cancer [44]. Studies have confirmed that activation of the Wnt signaling pathway is able to promote the epithelial-mesenchymal metastasis of prostate cancer, and induce the proliferation of prostate cancer cells [45, 46]. The present study has identified corroboratory evidence with respect to this pathway’s upstream miRNA and genes associated with prostatic cancer.

KRT5 is closely associated with prostate cancer-linked genes; furthermore, evidence was accumulated to suggest that KRT5 participates in the incidence of numerous types of cancer. For example, Breyer et al [41] demonstrated that there was a strong correlation of KRT5 expression with the recurrence and metastasis of nonmuscle invasive bladder cancer. Another study revealed that KRT5 could regulate the formation of cytoskeleton and venous invasion of cancer [42, 43]. Furthermore, Ricciardelli et al [29] verified the tumor suppressor function of KRT5 in reducing
Figure 12. Apoptosis of the PCSCs in each experimental group. A. Results of the flow cytometric analysis are shown. B. The apoptotic rate of the PCSCs after transfection for 48 h was determined. *P<0.05 vs. the negative control group; #P<0.05 vs. the miR-601 inhibitor + KRT5 group; &P<0.05 vs. the PRI-724 + KRT5 group. Data are expressed as the mean ± SD, and data were analyzed using one-way analysis of variance (ANOVA). The experiments were performed in triplicate. KRT5, keratin 5; PCSC, prostate cancer stem cell.
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Cancer stem cells, as the primary causative agents of tumor formation, resistance to chemoradiotherapy and cancer recurrence, share numerous features with normal stem cells, such as self-renewal and differentiation [47]. Accumulating evidence has demonstrated that miRNAs and genes exert regulatory roles in prostatic cancer by affecting PCSCs [48-50]. In order to investigate the effects of miR-601 and KRT5 on the biological characteristics of PCSCs, in the present study a series of experiments, including cell scratch healing, transwell, MTT assays and flow cytometric analyses, were performed. It was revealed that inhibition of miR-601 could activate KRT5 and inhibit activation of the Wnt signaling pathway, subsequently suppressing the migration, invasion and proliferation of the PCSCs, and their advancement. Nude mouse tumorigenicity assay is considered as a 'gold standard' for testing the tumorforming ability of tumor cells [51]. The results of our nude mouse tumorigenesis experiment revealed that the tumor volume in the miR-601 mimic group was much larger compared with that in the negative control group, although the tumor volume in the other groups was clearly decreased. Furthermore, miR-601 inhibition combined with KRT5 overexpression, as well as PRI-724 combined with KRT5 overexpression, led to a
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downregulation of the volume of the tumors compared with treated the mice with miR-601 inhibitor/KRT5/PRI-724. Therefore, it was possible to conclude that inhibition of miR-601 or overexpression of KRT5 could inhibit tumor growth by inhibiting activation of the Wnt signaling pathway. In addition, the immunohistochemical method was utilized to detect Ki-67 protein expression in each group’s lymph nodes, and the results demonstrated that inhibition of miR-601 or KRT5 overexpression was able to inhibit metastasis of the lymph node, and exert protective effects in prostate cancer progression.

In conclusion, the present study has demonstrated that downregulation of miR-601 decreases the rates of proliferation, invasion and metastasis of PCSCs by a mechanism that involves the interaction of miR-601 with KRT5, which subsequently influences the Wnt signaling pathway, highlighting the potential therapeutic value of miR-601 and KRT5 in treatment of prostatic cancer. The molecular mechanism of the present study is highlighted in Figure 15. However, the current study is still in the preclinical stages, and therefore numerous further studies are required in order that this avenue of research may provide proper direction for prospective therapies in the future.

Disclosure of conflict of interest

None.

Address correspondence to: Dr. Xinghuan Wang, Department of Urology, Zhongnan Hospital of Wuhan University, No. 169, Donghu Road, Wuchang District, Wuhan 430071, Hubei, P.R. China. Tel: +86-027-67812888; E-mail: wangxinghuan@whu.edu.cn

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