Extracellular matrix protein 1 (ECM1) is associated with carcinogenesis potential of human bladder cancer

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Introduction
Bladder cancer (BCa) is one of the most common urological malignant tumors worldwide, especially in elderly men, with almost 386,000 new cases and 150,000 deaths occurring annually. The incidence and mortality rate of BCa have the highest and the fifth highest among the urogenital carcinomas in China and the US, respectively. This is largely due to the inefficiency of current therapies for treating advanced BCa. Moreover, the molecular mechanism of BCa, involving the activation of proto-oncogenes and the inactivation of tumor suppressor genes in the carcinogenesis is far from clear, and the effective therapeutic targets are still lacking. Therefore, it is necessary to explore new therapeutic targets for BCa patients who are not sensitive to current therapy. Extracellular matrix protein 1 (ECM1) is a key player in multiple epithelial malignancies. However, the knowledge regarding the expression of ECM1 in BCa and the mechanisms by which ECM1 affects BCa tumor progression is unclear.

Materials and methods: ECM1 expression levels in BCa tissues and cells were detected by quantitative real-time PCR (qRT-PCR), immunohistochemistry and Western blot. ECM1 expression was suppressed by shRNAs. Cell Counting Kit-8 (CCK-8), luminescent cell viability assay and 5-ethyl-2′-deoxyuridine (EdU) assay were used to detect cell proliferation. Flow cytometry and transwell assay were used to evaluate cell apoptosis and invasion, respectively. All statistical analyses were performed by using the GraphPad Prism 7 software package.

Results: In this study, the expression of ECM1 in BCa specimens and cell lines was examined and displayed a significant increase compared with noncancerous counterparts, while ECM1-knockdown affected not only cell proliferation and migration, but also cell invasion ability and apoptosis potential, corresponding to the finding that ECM1 overexpression in BCa patients was associated with a poor prognosis. Additionally, after suppression of ECM1, the expression of glucose transporter 1 (GLUT1), lactate dehydrogenase (LDHA) and hypoxia-inducible factor 1α (HIF-1α), genes involved in Warburg effect regulation, were significantly decreased, and the lactate production was also obviously reduced in ECM1-silenced cells.

Conclusion: Our investigations revealed that the expression of ECM1 was closely associated with tumor cell growth, migration and apoptosis at least in part through regulation of Warburg effect, defining ECM1 as an effective predictor in the carcinogenesis and postoperative recurrence of human BCa.

Keywords: bladder cancer, extracellular matrix protein 1, proliferation, migration, invasion

Background: Bladder cancer (BCa) is a common urological malignant tumor worldwide, and recurrence and death still remain high. New therapeutic targets are needed to treat patients who are not sensitive to current therapy. Extracellular matrix protein 1 (ECM1) is a key player in multiple epithelial malignancies. However, the knowledge regarding the expression of ECM1 in BCa and the mechanisms by which ECM1 affects BCa tumor progression is unclear.
to current therapies, and elucidate the molecular mechanism of BCa carcinogenesis.\(^6\) \(^7\)

Extracellular matrix protein 1 (ECM1) is a glycoprotein expressed in epithelial organs and secreted into the extracellular stroma.\(^8\) Increased ECM1 expression has been reported in several malignancies, including invasive ductal breast carcinomas, laryngeal carcinomas, hepatocellular carcinomas, cholangio carcinomas, esophageal squamous cell carcinomas, gastric, colorectal, lung and thyroid cancers.\(^9\) \(^10\) Moreover, it is involved in various biological processes, such as cell proliferation, angiogenesis, migration and metastasis.\(^11\) \(^12\) ECM1 was found to promote metastatic progression and invasion in multiple cancer types, and overexpression of ECM1 has been incorporated into scoring models for indicating of poor clinical prognosis and metastatic potential in cancer patients.\(^13\) \(^14\) Compared with non-lymph node metastases, breast and lung tumors with lymph node metastases were more likely to be ECM1-positive.\(^15\) ECM1 overexpression in patients with invasive breast carcinomas confers a worse long-term prognosis.\(^16\) Further studies indicated that ECM1 induces tumor growth and metastasis by promoting angiogenesis and Warburg effect,\(^8\) or by enhancing the EGF signaling in breast cancer.

In the current study, we investigated the intracellular mechanisms by which ECM1 regulates metastatic behavior using aggressive BCa cell lines. Our results demonstrate that ECM1 was upregulated in BCa tissues and cell lines. Knockdown of ECM1 by shRNA inhibited cell proliferation, invasion and migration, while promoting the cell apoptosis, which might be mediated through attenuating the Warburg effect. Our findings define the essential role for ECM1 in the metastatic process and provide new insights into the role of ECM1 in BCa carcinogenesis.

### Materials and methods

#### Patients and specimens

A total of 17 fresh primary BCa tissues and noncancerous counterparts were collected from patients and quick-frozen in liquid nitrogen after radical cystectomy. None of the patients had received preoperative treatment and metastatic tumor specimens from other tissue origins were excluded. Written informed consents were obtained from all the patients or their relatives (with power of the attorney of the patient, who cannot read and write). The research protocol using human tissues was reviewed and approved by the Institutional Ethics Committee of the First Affiliated Hospital of Shenzhen University and the study was performed in accordance with the principles of the Declaration of Helsinki.

#### Cell culture and transfection

Human BCa cell lines (5637, BIU-87, T24 and SW780) and a normal urothelial cell line (SV-HUC-1) were purchased from the Institute of Cell Biology, Chinese Academy of Science (Shanghai, China). The cells were routinely grown as monolayers in phenol-free RPMI-1640 medium (5637 and BIU-87), DMEM (T24 and SW780), or F12K (SV-HUC1) (Gibco; Thermo Fisher Scientific, Waltham, MA, USA) supplemented with 10% charcoal-stripped FBS (Thermo Fisher Scientific), 100 U/mL streptomycin sulfate and 100 U/mL penicillin at 37°C in a humidified chamber containing 5% CO\(_2\). ECM1-silenced cells were generated by transfection with targeted ECM1 shRNA (shECM1: 5′-AACGAGGCC AGAGCAGTTTCAAGAGATGTGGAAGTGC TCTGCGCTTATTGTTGACC-3′) and nonspecific control shRNA (NC: 5′-TAATTGTCAATCAGAGTGC TTAAAGAGAAGACACTCTGATTGTACAAATTA-3′) using lipofectamine 3,000 transfection reagent (Thermo Fisher Scientific) according to manufacturer’s instructions.

#### cDNA preparation and qRT-PCR

The total RNAs from BCa tissues or cells were homogenized and isolated using the TRIzol reagent (Thermo Fisher Scientific) following the manufacturer’s instructions. Total RNA (1 µg) was converted to cDNA using a Revertra Ace qPCR RT Kit with gDNA Eraser (Toyobo Co Ltd, Japan). Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was used as the internal control. Other primers used were listed as follows: ECM1 forward: 5′-TCTCTTCAAGAGATGTGGAAGTGC TCTGCGCTTATTGTTGACC-3′, ECM1 reverse: 5′-AACGAGGAGAAGACACTCTGATTGTACAAATTA-3′, GLUT1 forward: 5′-ATGGGAAAGGTGAAAGTGC TCTGCGCTTATTGTTGACC-3′, GLUT1 reverse: 5′-TTCAAGAGAAGACACTCTGATTGTACAAATTA-3′, LDHA forward: 5′-CTCTTCAAGAGATGTGGAAGTGC TCTGCGCTTATTGTTGACC-3′, LDHA reverse: 5′-ATGGGAAAGGTGAAAGTGC TCTGCGCTTATTGTTGACC-3′, HIF-1α forward: 5′-ACCACTATGACCTGCTTTAGTGTGCG-3′, HIF-1α reverse: 5′-ATATCCAGGCTGACCTGCCC-3′.

Warburg effect associated genes: \(^9\) GLUT1 forward: 5′-AACGAGGAGAAGACACTCTGATTGTACAAATTA-3′, GLUT1 reverse: 5′-TCCTCCCATAGCAGGTGCC-3′, LDHA forward: 5′-ATGGGAAAGGTGAAAGTGC TCTGCGCTTATTGTTGACC-3′, LDHA reverse: 5′-ATGGGAAAGGTGAAAGTGC TCTGCGCTTATTGTTGACC-3′, HIF-1α forward: 5′-ACCACTATGACCTGCTTTAGTGTGCG-3′, HIF-1α reverse: 5′-ATATCCAGGCTGACCTGCCC-3′.

Quantitative real-time PCR (qRT-PCR) was performed in the ABI PRISM 7500 Fluorescent Quantitative PCR System (Thermo Fisher Scientific) by using SYBR Green Premix kit (Toyobo Co Ltd). The reaction system (15 µL) included 7.5 µL of PreMix, 1.5 µL of forward primer (2 µM), 1.5 µL of reverse primer (2 µM), 2 µL of cDNA diluted solution (10×) and 2.5 µL of deionized water. All the indicated samples were
normalized to GAPDH and then the relative expression levels were calculated using the 2^(-ΔΔ CT) formula.

**Cell proliferation assay**

Cell proliferation was measured by Cell Counting Kit-8 (CCK-8) assay, luminescent cell viability assay, and 5-ethyl-2′-deoxyuridine (EdU) labeling assay. In brief, 1×10⁴ cells were seeded in a 96-well plate. 100 µL fresh medium with 10% CCK-8 reagents (TransGen Biotech Co., Ltd, Beijing, China) was replaced into each well at 0, 24, 48, and 72 hours and incubated for another 2 hours at 37°C. The absorbance at 450 nm was detected by using an ELISA microplate reader (Bio-Rad Laboratories Inc., Hercules, CA, USA). For luminescent cell viability assay, 1×10⁴ cells suspended with 100 µL fresh media were seeded into 96-well plate. 72 hours later, 100 µL luminescent cell viability assay reagents were added into each well and incubated for 10 minutes. The luminescent signals were recorded in a Glo Max Discover system (Promega Corporation, Fitchburg, WI, USA). To directly observe the cell proliferation, EdU incorporation experiments were conducted according to the manufacturer’s specifications. Briefly, 48 hours after transfection, the EdU reagents (RIBOBIO, Guangzhou, China) were added into each well in a final concentration of 50 µM. Two hours later, cells were fixed with 4% paraformaldehyde in PBS at room temperature. Before 1× Apollo solution incubation for half an hour at room temperature in dark, cells were washed three times in PBST (PBS containing 0.1% Triton X-100). After that, cells were subjected to 1× Hoechst nuclear dye staining for 30 minutes and then observed under the fluorescence microscopy (DM IL LED, Leica Microsystems, Wetzlar, Germany). Each experiment was repeated at least three times.

**Cell migration assays**

Cell migration was evaluated in the ECM1-silenced 5637 and BIU-87 cell lines with the wound healing assay. Cells were seeded in a 6-well plate and grown to confluence (1×10⁵ per well) in 1,640 medium for 24 hours. The monolayer cells were scratched with a sterile 200 µL pipette tip to create clear lines. Cells were washed twice with serum-free 1,640 medium to remove floating cells and then cells were continuously cultured in the medium with 1% FBS for 24 hours. Migrating cells from the edge of the wound were recorded at 0 and 24 hours using an Olympus IX51 inverted fluorescence microscope and digital camera system (Olympus Corporation, Tokyo, Japan). The cell migration was evaluated by comparing the remaining cell-free area with the initial scrape line and calculated using ImageJ software (http://rsbweb.nih.gov/ij; NIH, Bethesda, MD, USA).

**Invasion assay**

Transwell assay was conducted to assess cell invasion using 24-well Matrigel pre-coated cell culture inserts (BD Biosciences, San Jose, CA, USA). Briefly, cells were detached from the culture plates, washed and re-suspended. 200 µL aliquot (1×10⁵) of cells in 1,640 medium (1% FBS) was plated into the upper chamber of each insert, and filled the lower chamber with 500 µL complete medium. After 24 hours of cultivation, noninvasive cells were removed from the upper surface of the insert using a cotton swab, whereas invaded cells on the bottom layer surface were rinsed in 1× PBS, fixed with methanol for 20 minutes, stained with 0.1% crystal violet dye (Sigma-Aldrich Co., St Louis, MO, USA) for 25 minutes, followed by washing three times with 1× PBS. The invaded cells were then observed under an inverted light microscope and photographed after drying. Finally, the chamber inserts with invaded cells were soaked into 1 mL of 33% acetic acid for 10 minutes to wash out the crystal violet. The absorbance at 570 nm was measured using a microplate reader (Bio-Rad Laboratories Inc.). Experiments were performed in triplicate.

**Flow cytometry analysis of cell apoptosis**

Cell apoptosis was determined using flow cytometry assay. ECM1-silenced cells were harvested and washed with ice-cooled 1× PBS. The cells were then co-mixed with 5 µL annexin V-FITC and 5 µL propidium iodide (TransGene, Beijing, China) in 100 µL annexin V binding buffer and incubated in dark at room temperature for 15 minutes according to the manufacturer’s specifications. The proportion of apoptotic cells was analyzed by flow cytometry (EPICS, XL-4, Beckman Coulter, Inc., Brea, CA, USA). The experiments were performed in triplicate.

**Lactate assay**

The levels of lactate production were measured with a lactate assay kit (Sigma-Aldrich Co.). 2×10⁵ cells were plated and transfected in 10-cm culture dishes and grown for 24 hours. FBS-free media was replaced into each culture dish for a further incubation of 8 hours. Lactate production was measured with culture media collected from each sample according to the manufacturer’s protocol and the optical density was recorded at 570 nm.
Western blot analysis
Cells were transfected and grown in 6-well plates, harvested and lysed with lysis buffer containing 10 mM Tris-HCl, 5 mM EDTA, 50 mM NaCl, 50 mM sodium fluoride, 30 mM sodium pyrophosphate, 1% Triton-X, 200 µM sodium orthovanadate, 1 mM phenylmethysulfonyl fluoride, 1 µg/mL peptatin, 2 µg/mL leupeptin and 5 µg/mL aprotinin. Protein concentration in the cell lysates was determined using the BCA protein assay kit (Thermo Fisher Scientific). Equal amounts of whole protein extracts were separated on 10% SDS-PAGE gels for detecting expression of ECM1 and internal control β-actin, respectively. After transferred to polyvinyl difluoride membranes, samples were blocked in 5% skimmed milk in TBST for 1 hour and then incubated overnight with the specific antibodies (Abnova Corporation, Taipei City, Taiwan). The levels of proteins were detected with horseradish peroxidase-linked secondary antibodies and visualized using the ECL® System (GE Healthcare, Milan, Italy).

Immunohistochemical staining
Tissue specimens were fixed in 10% neutral formalin, embedded in paraffin, and cut into 4-µm sections. The sections were deparaffinized with xylene and alcohol, processed with antigen retrieval by being boiled in citrate buffer (0.01 M, PH 6.0) for 20 minutes and then cooled down at room temperature. After that, the slides were incubated in a 10% hydrogen peroxide solution for 15 minutes and washed with PBS for 5 minutes. The slides were then immunostained with primary antibody of rabbit anti-ECM1 (1:1,000) antibody (Abnova Corporation) at 4°C overnight. After rinsing with PBS for 5 minutes, the sections were incubated with FITC-conjugated secondary antibody and DAPI (Thermo Fisher Scientific) at 4°C for 30 minutes. The slides were then washed with PBS for three times, the signal was detected under a fluorescent microscope (DM IL LED, Leica Microsystems).

Statistical analyses
The results reported were obtained from at least three independent experiments and the data were expressed as mean ± SD. Statistical tests were performed by Student’s t-test using the GraphPad Prism 7 software package (GraphPad Software, Inc., La Jolla, CA, USA) (www.graphpad.com/) and the significant differences were considered at P<0.05, P<0.01, P<0.001.

Results
The ECM1 level was upregulated in BCa
To explore the effects of ECM1 on BCa development, we first examined the relative expression levels in BCa tissues (from 17 BCa patients, characterized in Table 1) and 2 cell lines by using qRT-PCR and Western blot. The ECM1 levels were separately normalized to paired-matched adjacent noncancerous tissues or control cell lines. The results showed that the expression of ECM1 was upregulated in 82.3% of cancer tissues (14/17) compared with noncancerous counterparts (Figure 1A). We further confirmed the expression pattern by Western blot (Figure 1B) and immunohistochemical staining (Figure 1C) in one patient with low ECM1 mRNA expression and 2 patients with higher mRNA expression of ECM1 as representatives. ECM1 was mainly expressed in the cytoplasm of BCa cells (Figure 1C). In addition, RNA expression data from The Cancer Genome Atlas (TCGA) indicated that patients with higher levels of ECM1 had poorer survival than those with lower expression by using the Kaplan–Meier method (Figure 1D), and that increased ECM1 levels were highly correlated with BCa clinical pathological grading (Figure 1E). We next verified that the expression of ECM1 was increased in 5637 cells and BIU-87 cells in contrast to the normal bladder cell line SV-HUC1 (Figure 1F and G). Together, these results indicate that the expression of ECM1 is significantly elevated in BCa and that ECM1 may play an oncogenic role in BCa progression.

Knockdown of ECM1 suppressed BCa cell growth
The functional role of ECM1 on cell proliferation in BCa was tested by CCK-8, luminescent cell viability assay, and EdU incorporation assays. The relative expression levels of ECM1 were obviously decreased in 5637, BIU-87, T24 and SW780 cells by shRNA (Figure 2A and E). The CCK-8 assay and luminescent cell viability assay showed that knockdown of ECM1 significantly suppressed cell proliferation compared with negative controls (Figure 2B, F–I). The EdU assay was conducted to further detect function of ECM1 in promoting cell proliferation. As shown in Figure 2C, in contrast to the NC group, the rate of EdU positive cells was significantly
decreased in the shECM1 group. The quantitative results proved that the number of EdU positive cells in the shECM1 group was obviously reduced in 5637 and BIU-87 cells (Figure 2D). These data further support the notion that ECM1 may promote cell proliferation in BCa.

Knockdown of ECM1 inhibited cell migration and invasion

The impacts of ECM1 on BCa cell migration were investigated by wound-healing assay. The results showed that the migration rates of the cells with silenced-ECM1 were obviously
Figure 2 Evaluation of growth and proliferation in ECM1-silenced BCA cells.

Notes: (A) Downregulation of ECM1 by shRNAs in BCA cell lines 5637 and BIU-87. (B) Cell proliferation assay by CCK8 method. (C) Cell proliferation assay by EdU incorporation (magnification ×100). (D) The quantification of (C). (E) ECM1 protein expression in BCA cell lines transfected with shECM1 or NC. (F–I) Cell proliferation assay by luminescent cell viability assay. Data are shown as mean ± SD (*P<0.05, **P<0.01, ***P<0.001).

Abbreviations: NC, negative control; ECM1, extracellular matrix protein 1; EdU, 5-ethyl-2'-deoxyuridine; RLU, relative light unit; BCAs, bladder cancer; CCK8, Cell Counting Kit-8.
repressed compared with those NC groups (Figure 3A and C). The quantification of cell migration showed a significant diminution in cells treated with $ECM1$ shRNA compared with negative controls (Figure 3B and D). Subsequently, we assessed the effect of $ECM1$ on cell invasion with transwell assay. As shown in Figure 3E–G, compared with the control group, knockdown of $ECM1$ significantly decreased the number of invading cells. These results collectively suggest that $ECM1$ silencing may affect the metastatic capacity of both cell lines by inhibiting cell migration and invasion.

**Knockdown of $ECM1$ changed cell apoptosis and gene expression**

Flow cytometry assay was performed to detect the cell apoptotic changes caused by knockdown of $ECM1$ in BCa cells. Our results showed that there was a higher ratio of apoptotic cells in $ECM1$-silenced 5637 and BIU-87 cells compared with those of negative control cells (Figure 4A and C). The quantitative results also confirmed these results (Figure 4B and D). The Warburg effect is an oncogenic metabolic switch that is commonly observed in cancer cells. We demonstrated that silencing of $ECM1$ reduced the expression of genes involved in Warburg effect regulation, including $GLUT1$, $LDHA$, and $HIF-1\alpha$ (Figure 4E and F). Consistent with this result, we detected lower levels of lactate production in $ECM1$-silenced cells, further confirming the involvement of Warburg effects (Figure 4G and H). Together, these findings suggest that loss-of-function of $ECM1$ induced apoptosis of BCa and the Warburg effect might be involved in the development of BCa.

**Discussion**

BCa is one of the most common malignancies worldwide, imposing heavy economic burdens on both the society and individuals. Our study demonstrated that $ECM1$ was highly expressed in BCa tissues, cell lines, and highly correlated with the histological grades and poor survival in patients with BCa. Knockdown of $ECM1$ in
BCa cells affected not only several cellular characteristics, but also affected the expression of genes involved in the Warburg effect. As such, we propose that ECM1 is a key regulator in the occurrence and progression of BCa. ECM1 overexpression has been correlated with a poor prognosis in patients with breast cancer\textsuperscript{11,21} and hepatocellular cancer.\textsuperscript{22} This is consistent with the elevated expression of ECM1 in BCa specimens compared to the normal counterparts from the same patients, suggesting its role in carcinogenesis and invasiveness of BCa cells.

**Figure 4** (Continued)
It has been speculated that ECM1 as a “biological glue” binding to components of the dermal-epidermal junction in the framework of normal skin. Tumor cell invasion needs several steps, such as extracellular matrix adhesion and degradation, cell migration and detachment. Altered expression of this “glue” in tumor cells might alter attachment and invasion, and subsequently affect metastasis and prognosis. Indeed, our results demonstrated that knockdown of ECM1 by shRNA in 5637 and BIU-87 cells resulted in alterations in cell proliferation, migration, invasion and apoptosis. Similar results have been reported in cholangiocarcinoma cell lines with ECM1 silenced.

Because that one gene is unlikely to be the sole effector in carcinogenesis, we noted that ECM1 does indeed appear to regulate the expression of genes known to be important in Warburg effect regulation. Knockdown of ECM1 reduced cell proliferation and invasiveness by inhibiting the expression of GLUT1, LDHA and HIF-1α. Similar results have been observed in breast cancer cell line BT-474. The Warburg effect mediates a metabolic switch for ATP production from oxidative phosphorylation to glycolysis, converting most of the incoming glucose to lactate. Just as we expected, the lactate production was significantly decreased in ECM1-silenced cells, confirming the involvement of Warburg effect in the development of BCa.

**Conclusion**

ECM1 was highly expressed in BCa tissues and cell lines. Downregulation of ECM1 inhibited cell proliferation, migration and invasion by regulating the Warburg effect related genes. Therefore, our findings provide novel insights into the role of ECM1 in BCa development.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**