Fucosyltransferase VII promotes proliferation via the EGFR/AKT/mTOR pathway in A549 cells

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Abstract: Fucosyltransferase VII (FUT7) is one of a1,3-fucosyltransferases family that catalyzes the final fucosylation step in the synthesis of Lewis antigens and generates a unique glycosylated product sialyl Lewis X (sLe\(^X\)). sLe\(^X\) can serve as ligands for E- or P-selectin expressed on the cell surface and results in cancer metastasis and angiogenesis. However, the molecular biological mechanisms of FUT7 elevation in neoplastic cells are still largely unknown. In this study, we examined the impact of FUT7 on cell proliferation and migration in A549 cells by colony formation assay, cell cycle assay, gelatin zymography, wound-healing assay, transwell invasion assay and Western blot. In addition, we identified that FUT7 activated EGFR/AKT/mTOR signal pathway that correlated with sLe\(^X\) augmentation. In conclusion, FUT7 overexpression augments sLe\(^X\) synthesis to trigger cell proliferation via the activation of EGFR/AKT/mTOR signaling pathway, which indicated that FUT7 may be a potential therapeutic target for epithelial cancers with a high expression of FUT7 and sLe\(^X\).

Keywords: fucosyltransferase VII, epidermal growth factor receptor, lung cancer, proliferation, signal pathway

Introduction

Lung cancer is a high malignant carcinoma that has been reported to be the leading cause of cancer death in the US.¹ Despite the advanced diagnostic techniques for early detection of lung cancer, the prognosis of lung cancer in patients is still unsatisfactory. The mortality remains high even among patients with early-stage lung cancer.² Because of the limited knowledge of lung cancer, the therapeutic technology still needs further development. Therefore, the identification of molecules related to the progression of lung cancer will facilitate the development of therapeutic strategies for lung cancer.

It has been reported that overexpression or mutation of EGFR induced excessive proliferation of cells in lung cancer.³,⁴ Activation of EGFR signaling promotes cell proliferation, migration and adhesion via stimulating the downstream signal pathways, such as RAS/BRAF/MEK/ERK, PI3K/AKT/mTOR, JAK/STAT and PLC/PKC.⁵,⁶ EGF and TGF-α secreted by tumor cells combining with abnormal expression of EGFR activate the downstream signaling pathway that results in tumor malignant behavior and poor prognosis.⁷ In view of the important role of EGFR in the progression of lung cancer, the targeted therapy has become a new therapeutic method and achieved positive outcomes. EGFR-targeting drugs are widely used for clinical application, yet with unsatisfactory therapeutic effect. Therefore, searching for an effective EGFR-sensitizing target may enhance the therapeutic effect of EGFR-targeted therapy and improve the prognosis of lung cancer patients.
Glycosylation is a common post-translational regulator of eukaryotic gene expression. The synthesis of carbohydrate antigens is catalyzed by glycosyltransferases. a1,3-Fucosyltransferases (a1,3-FUTs) catalyze the final fucosylation step in the synthesis of Lewis antigens, which are a series of sugar chains derived from Galβ1-3/1-4GlcNac backbone. Six members of this family (FUT3, 4, 5, 6, 7 and 9) have already been cloned and identified. Each enzyme has a specific acceptor-substrate binding pattern and generates a unique range of glycosylated products. Lewis antigens are mainly located at the outer chains of glycolipids and the O- and N-linked sugar chains on the cell surface of glycoproteins. Fucosyltransferase VII (FUT7) catalyzes the synthesis of sialyl Lewis X (sLeX; NeuNAcα2-3Galβ1-4[Fucα1-3]GlcNAc), which is its only product. sLeX can serve as ligand for E- or P-selectin expressed on the cell surface and mediates the adhesion of malignant cells that result in cancer metastasis and angiogenesis.

Ogawa et al first showed that FUT7 related to metastasis and remained a statistically significant prognostic factor in lung cancer. In early meta-analysis, we showed that sLeX overexpression was associated with tumor metastasis, recurrence and overall survival in cancer patients and played an important role in cancer prognosis. Farhan et al have shown that anti-Lewis Y antibody inhibited the EGF-stimulated EGFR signaling in A431 cells and prevented cell proliferation. Downregulation of FUT3 and FUT6 caused inhibition of colon carcinoma cell proliferation. However, the regulation of FUT7 expression and its function in the proliferation of non-small lung cancer cells are largely unknown. The present study further identifies the correlation between FUT7 and EGFR in lung cancer cells.

**Materials and methods**

**Reagents**

Kinase inhibitor LY294002 was purchased from Beyotime Institute of Biotechnology (Shanghai, China). CCK-8 kit was purchased from Dojindo Molecular Technologies (Kumamoto, Japan). FUT7 expression construct and the blank vector were designed and synthesized by GenePharma (Shanghai, China). anti-AKT, anti-EGFR, anti-mTOR and anti-PCNA antibodies were purchased from Proteintech (Wuhan, China); anti-pEGFR, anti-pAKT (ser473) and anti-pmTOR (ser2448) antibodies were purchased from Bioworld Technology, Inc (St Louis Park, MN, USA); anti-FUT7 monoclonal antibody was purchased from Sigma-Aldrich Co. (St Louis, MO, USA) and anti-sLeX antibody was purchased from Merck Millipore (Billerica, MA, USA).

**Cell culture and transfection**

FUT7 overexpression plasmid pIRE2-EGFP-FUT7 (GenePharma) was constructed following the manufacturer’s instructions. The human lung cancer cell line (A549) was purchased from the Type Culture Collection of the Chinese Academy of Sciences (Shanghai, China). A549 cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM; Hyclone, Logan, Utah, USA) supplemented with 10% fetal bovine serum, 100 IU/mL penicillin and 100 μg/mL streptomycin at 37°C with 5% CO2. pIRE2-EGFP-FUT7 was transiently transfected into A549 cells using 0.1 μg (low dose, FUT7 cDNA-1) or 0.4 μg (high dose, FUT7 cDNA-2) of plasmids in the presence of 2 μL of Lipofectamine reagent per manufacturer’s instructions.

**Colony formation assay**

Cells with different treatments were seeded in six-well plates at an initial cell density of 200 cells/well. Colonies were allowed to form for 2 weeks and stained with 0.05% crystal violet after methanol fixation.

**Analysis of cell cycle**

Cells were cultured in six-well plates and allowed to grow to 75%–80% confluency. Nonadherent cells were removed by gentle washing with PBS. Cells were trypsinized, collected and washed twice with PBS. Cell pellets were resuspended in 0.5 mL of PBS and fixed in 4.5 mL of 70% ethanol overnight. Cells were collected by centrifugation, and the pellets were resuspended in 0.2 mg/mL of propidium iodide (PI) containing 0.1% Triton X-100 and 0.1 mg/mL RNase A. The cell suspension was incubated in the dark for 30 min at room temperature and subsequently analyzed using FACScan flow cytometer for DNA content. The percentage of cells in different phases of the cell cycle was sorted using a ModFit 5.2 computer program.

**Western blot**

Cells were lysed using radioimmunoprecipitation assay (RIPA) buffer (25 mM Tris-Hcl [pH 7.6], 150 mM NaCl, 1% NP-40, 1% sodium deoxycholate, 0.1% sodium dodecyl sulfate [SDS]) with protease inhibitors (EMD Biosciences, San Diego, CA, USA). After sonication for 2 min at 4°C and centrifugation for 10 min at 4°C, the supernatant was taken as the total cell lysate. Protein concentration was quantified using the Bradford method. Equal amounts of total protein (50 μg) were analyzed by SDS polyacrylamide gel electrophoresis (SDS-PAGE). Following transfer to the nitrocellulose (NC) membrane, the blot was probed with...
primary antibodies as indicated and then incubated with horseradish peroxidase (HRP)-labeled secondary antibodies for visualization using enhanced chemiluminescence reagents (Thermo Fisher Scientific, Waltham, MA, USA). The images were obtained by the Bio-Rad Imaging System.

**Gelatin zymography**
Cells were grown in six-well plates. The supernatants after the different treatments were collected. The supernatant protein (40 μg/lane) was loaded and electrophoresed on 8% SDS-PAGE copolymerized with 1% gelatin. After electrophoresis, the gels were washed five times in 2.5% Triton X-100 (20 min each) and two times in buffer without Triton X-100 to remove Triton X-100 and then incubated in 50 mmol/L Tris-Cl (pH 7.6) and 5 mmol/L CaCl₂ (18 h, 37°C). The gels were stained with 0.1% Coomassie Blue R-250 and destained in 10% isopropanol and 10% acetic acid in H₂O. MMP2 and MMP9 levels were detected as transparent bands on the blue background of a Coomassie Blue-stained gel.

**Immunoprecipitation**
Cells were lysed in RIPA buffer (Beyotime Institute of Biotechnology), and 500 μg of protein was incubated with an anti-EGFR antibody for 1 h at 4°C. Antibodies were precipitated using Protein G-coated Dynabeads (Thermo Fisher Scientific), and bound protein was eluted in Laemmli buffer (Thermo Fisher Scientific) for 10 min at 70°C.

**Wound-healing assay**
Cells were seeded in six-well plates for 24 h. The monolayers were scratched with a 200 μL pipette tip and washed with media without serum to remove the detached cells. The wounded areas were observed and imaged using a microscope.

**Transwell invasion assay**
For invasion assays, cells (1×10⁵/well) were placed into the upper chamber with the Matrigel-coated membrane, which was diluted with a serum-free culture medium. Cells were suspended in 200 μL of serum-free Roswell Park Memorial Institute (RPMI)-1640 medium and seeded into the upper chamber. In the lower chamber, 800 μL of the RPMI-1640 medium supplemented with 10% fetal bovine serum was added. After incubation for 24 h at 37°C and 5% CO₂, the chambers were fixed with 100% methanol for 20 min and stained with 0.1% crystal violet for 15 min. Images were captured with an Olympus BX83 fluorescence microscope (Olympus Corporation, Tokyo, Japan).

**Cell viability assay**
Cell proliferation inhibition was evaluated using the CCK-8 (Dojindo Laboratories, Kumamoto, Japan) according to the manufacturer’s instructions. A549 cells (6×10⁴/90 μL/well) were plated into 96-well plates in triplicate and cultured for 24 h before the onset of treatment. Then, cells were treated with LY294002 (10, 20, 30, 40, 50 and 60 μM) for 24 h in 96-well plates with 2,000 cells/well. Then, 20 μL of CCK-8 reagent was added to each well for 1 h, followed by light absorbance measurement at a wavelength of 450 nm.

**Immunocytochemistry**
Cells were seeded on glass coverslips and then fixed with PBS containing 4% formaldehyde for 10 min and permeabilized with 0.2% Triton X-100 in PBS for 10 min at room temperature. After washing with PBS, endogenous peroxidase activity was suppressed by 3% H₂O₂ and blocked with goat serum or 5% bovine serum albumin (BSA). Diluted primary monoclonal rabbit antibodies against sLeα (dilute 1:100) were added and incubated at 4°C overnight. As secondary reagents, biotin-labeled anti-IgG and an avidin–biotin HRP complex were used Zhongshan jinqiao (Beijing, China) followed by staining with chromogen diaminobenzidine (Zhongshan) until a brown color developed. Slides were counterstained with Mayer’s hematoxylin and differentiated in a solution containing 1% hydrochloric acid and 99% ethanol. Sections were dehydrated, and a transparent cover slip was added to enable observation by microscopy. All the immunohistochemical photographs were analyzed using Image-Pro Plus (IPP; Media Cybernetics, Inc., Rockville, MD, USA).

**Statistical analysis**
Results are expressed as the mean ± standard error of the mean (SEM) of at least three independent experiments. The statistical significance of differences among groups was analyzed by one-way analysis of variance (ANOVA) and the Student–Newman–Keuls (SNK) q test using SPSS 17.0 software (SPSS Inc., Chicago, IL, USA). Differences were considered to indicate a statistically significant result with a P-value <0.05.

**Results**
**FUT7 induces sLeα expression in A549 cells**
To examine the relationship between FUT7 and sLeα, two FUT7 cDNA constructs (cDNA-1 and cDNA-2) and the blank vector as control transfected A549 cells. As shown in Figure 1A, when FUT7 was overexpressed by FUT7
cDNA constructs, FUT7 levels were increased, concomitant with a notable increase in sLeX, indicating that FUT7 was a factor for promoting sLeX expression. In Figure 1B, we used immunocytochemistry to detect sLeX expression in different FUT7 expression A549 cells. It showed similar result as Western blot.

Overexpression of FUT7 promotes cell growth

We measured cell proliferation utilizing colony formation assay (Figure 2A) and found that overexpression of FUT7 significantly increased the number of formed colonies. The impact of FUT7 overexpression on cell proliferation was further determined by cell cycle analysis (Figure 2B). Flow cytometry with PI-stained cells showed that A549-vector control cells were presented in G0/G1 (82.46%±1.89%), S (17.01%±2.04%) and G2/M (0.53%±0.01%) phases. While in A549-FUT7 cDNA cells, S phase fraction was increased (36.62%±1.48%, P<0.01) and G0/G1 fraction was decreased (0.00%±0.00% for A549-FUT7 cDNA cells, P<0.01). To further evaluate the impact of FUT7 overexpression on cell proliferation, PCNA expression was detected by Western blot (Figure 2C). A significant increase in PCNA expression was observed in A549-FUT7 cDNA cells.

Overexpression of FUT7 promotes cell migration and invasion in A549 cells

We observed the migratory capacity of the cells through wound healing assay or scratch assay (Figure 3A). We found that the scratch was almost empty after 24 h in the A549-vector group, while the cells in the A549-FUT7 cDNA group showed higher migratory ability, as they filled over...
half of the scratch within 24 h. As the wound healing assay provides only semi-quantitative results, we proceeded with a transwell migration assay to evaluate the cell migratory capacity more precisely. The cells in the A549-FUT7 group showed a significant higher migratory ability compared with the cells in the A549-vector group (P<0.05; Figure 3B). We then emphasized the result by using the gelatin zymography assay and found that the activities of MMP2 and MMP9 were both increased in the A549-FUT7 cDNA group (Figure 3C).

**FUT7 promotes proliferation via the EGFR/AKT/mTOR pathway in A549 cells by inducing sLe\textsuperscript{X} generation**

To elucidate the molecular mechanism underlying the augmented proliferation by FUT7, the EGFR/AKT/mTOR pathway was specifically analyzed in the present study. Using gene overexpression combined with anti-sLe\textsuperscript{X} antibody blocking, it was found that FUT7 overexpression increased the activity of EGFR, AKT and mTOR as reflected by the phosphorylated form of EGFR, AKT and mTOR (Figure 4A). In addition, anti-sLe\textsuperscript{X} antibody inhibited the activation of EGFR/AKT/mTOR signaling pathway. In addition, we used immunoprecipitation to detect the interaction between sLe\textsuperscript{X} and EGFR; it was found that there is sLe\textsuperscript{X} modification on EGFR, and the expression increased in FUT7 overexpression cells (Figure 4B). To further evaluate the involvement of EGFR/AKT/mTOR signaling pathway in FUT7 overexpression-induced activation on cell proliferation, we used LY294002 to inhibit AKT activation. To determine the working concentration of LY294002, CCK-8 assay was performed to evaluate the inhibitory rate of LY294002 on A549 cells. As shown in Figure 5A, A549 cells with LY294002 treatment displayed an increased inhibitory rate; the inhibition concentration of 10% of LY294002 against A549 cells.
cells was 10±0.29 μM. Our result showed that inactivation of AKT signaling significantly inhibits the cell proliferation induced by FUT7 overexpression. Increases in PCNA expression was seen in A549-FUT7 cDNA cells, while the inhibited expression of PCNA was observed in A549-FUT7 cDNA cells combined with LY294002 treatment (Figure 5B). The number of colonies in A549-FUT7 cDNA cells combined with LY294002 treatment was significantly decreased compared with those in A549-FUT7 cDNA cells (Figure 5C). The Akt activity was assessed by Western blot with a phospho-specific antibody against phosphorylated Ser473, which has been reported to be constitutively active in A549 cells under normal growth conditions. We found that the LY294002 treatment combined with FUT7 overexpression led to a marked decrease in the level of phosphorylated Akt compared with that of the A549-FUT7 cDNA cells (Figure 5B).

**Discussion**

The carbohydrate moieties of cell surface glycoconjugates have been shown to play important roles in many biological processes. The multiple linkages, branching patterns and terminal modifications in these glycans are the main causes for their functional diversity. Over the last decades, important discoveries have been made in the field of glycoscience. It has been well studied that the neoplastic processes result in fundamental changes in the glycosylation patterns of cell surface glycoconjugates. Lewis antigens are a group of oligosaccharide residues constituting the end of various carbohydrate chains of glycoproteins and glycolipids protruding from the outer surface of mammalian cell membranes. Sialylated Lewis antigens sLeX are potentially involved in many cell adhesion events because they act as specific E-selectin ligands. The alteration in cell surface sLeX oligosaccharide expression has been reported to be correlated with many kinds of cancers. Our previous meta-analysis study has shown that sLeX overexpression is associated with tumor metastasis, recurrence and overall survival in cancer patients and plays an important role in cancer prognosis. Komatsu et al reported that serum sLeX concentration was an independent predictive factor for skip metastasis in non-small cell lung cancer (NSCLC). Therefore, exploring the mechanism of sLeX antigen synthesis is important for developing effective sLeX targeting therapeutic strategy. Biosynthesis of sLeX requires addition of fucose to N-acetyllactosamine by enzymatic catalysis. Among these enzymes, FUT7 was thought to participate in the biosynthesis of sLeX. In the present study, we established FUT7-overexpressed cell
colonies. We found that FUT7 overexpression significantly augmented the sLe\(^\alpha\) antigen synthesis compared with that of the mock control cells (Figure 1).

It has been reported that abnormal expression of FUT7 is closely related to poor prognosis in lung cancer.\(^1\) In the study of Wang et al,\(^2\) the growth of hepatocarcinoma cells was significantly stimulated via the cyclin-dependent kinase inhibitor p27Kip1 by transfecting of FUT7 cDNA. Laubli et al\(^3\) have found that deficiency of induced endogenous L-selectin ligands caused by a low expression of FUT7 led to a low incidence of metastasis of hematogenous carcinoma in mice. We evaluated the effect of FUT7 expression on cell growth and migration using human adenocarcinoma A549 cells. We found that FUT7 overexpression significantly increased the cell growth. Cellular proliferation was detected by using flow cytometry. FUT7 overexpression enhanced the DNA replication through the cell cycle arrest at the S phase (Figure 2). It has been shown that overexpression of FUT7 enhanced the colorectal carcinoma metastases via promoting the carbohydration of glycoprotein CD24.\(^4\) In the present study, we represented that migration and invasion capability of A549 cells were enhanced after the transfection of FUT7 cDNA (Figure 3). These results suggested that FUT7 plays a critical role in cell growth and in its capability of migration and invasion.

The underlying mechanism of sLe\(^\alpha\) regulating tumor progression still needs further exploration. It has been studied that overexpression of FUT7 led to activation of the InR pathway, which is due to the high expression of sLe\(^\alpha\) on the glycans of cell-surface receptors.\(^5\) Liu et al\(^6\) revealed that using FUT7 antisense oligodeoxynucleotide leads to reduced expression of both FUT7 and sLe\(^\alpha\) and thereby inhibited the adhesion of embryo cells to endometrium. Using in vivo models, the adhesion rate was significantly reduced when the sLe\(^\alpha\) antigen on the surface of endometrial cells was blocked by the anti-sLe\(^\alpha\) antibody.\(^7\) The dysregulation of EGFR signaling has been well documented to contribute to the progression of NSCLC. The AKT signaling pathways are downstream of EGFR and deregulated via genetic and epigenetic mechanisms in many human cancers and play a pivotal role in oncogenesis.\(^8\) It has been revealed that anti-Lewis Y antibody (Lewis Y antigen is catalyzed by fut4, a member of fucosyltransferase gene family) effectively blocked the EGFR-stimulated phosphorylation of MAPK through inactivating the EGFR signaling in A431 cells and prevented cell proliferation.\(^9\) Based on these studies, we hypothesized that blocking the sLe\(^\alpha\) antigen with anti-sLe\(^\alpha\) antibody may inhibit the EGFR/Akt/mTOR signaling pathway. In the present study, we verified our hypothesis by using PI3K/Akt inhibitor LY294002 and anti-sLe\(^\alpha\) antibody. Meantime, the immunoprecipitation result showed that EGFR glycoprotein is modified by sLe\(^\alpha\) that may participate in the phosphorylation of EGFR (Figures 4 and 5), but the underlying mechanism between the sLe\(^\alpha\) modification of EGFR and EGFR activation still needs further study. Our results indicated that anti-sLe\(^\alpha\) antibody inhibited tumorigenesis via inactivation of the EGFR/Akt/mTOR signaling pathway. FUT7 overexpression promotes tumorigenesis in NSCLC through activation EGFR/Akt/mTOR signaling pathway by augmenting the sLe\(^\alpha\) synthesis.

**Conclusion**

FUT7 played an important role in cell growth and proliferation via promoting the sLe\(^\alpha\) antigen synthesis and EGFR/AKT/mTOR signal pathway activation in the A549 cell line, which indicated that FUT7 may be a potential therapeutic target for epithelial cancers with a high expression of FUT7 and sLe\(^\alpha\).

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**Disclosure**

The authors report no conflicts of interest in this work.

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