Alpha-(1,6)-fucosyltransferase (FUT8) affects the survival strategy of osteosarcoma by remodeling TNF/NF-κB2 signaling

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INTRODUCTION

Osteosarcoma (OS) is the most frequent and aggressive primary malignant neoplasm diagnosed in the skeletal system [1]. Many patients have to face death threat after diagnosis due to its rapid progression, tendency to metastasize to the lungs, and high recurrence rate [2]. Since the advent of a comprehensive therapeutic strategy in the 1970s, which includes neoadjuvant chemotherapy, surgery, and radiotherapy, the 5-year overall survival rate increased for localized OS [1]. However, the survival rate has remained unchanged for decades, leaving 30–40% of patients who still cannot benefit from comprehensive therapy [3]. Therefore, it is important to identify all the factors that contribute to OS progression. Identification of such factors could pave the way for the development of novel clinical treatments for OS.

Glycosylation is one of the most important post-translational modifications of proteins in all cells [4], as it plays an indispensable role in maintaining many biological functions of proteins. Aberrant glycosylation is closely linked to a variety of diseases, especially cancers, and is due to alternations in the expression of glycosyltransferases [5]. Aberrant glycosylation affects well-known intrinsic traits of tumors, such as proliferation, metastasis, angiogenesis, and drug resistance, among others [6–9]. Fucosylation is a significant kind of glycosylation that is catalyzed by fucosyltransferases (FUTs) [10]. FUTs can be divided into four types according to their catalytic products: FUT1-2, FUT3-7, FUT9-11, and FUT8 [11, 12]. FUT8 exclusively catalyzes the formation of α-1,6 fucosylation—also known as core fucosylation—by connecting fucose to the inner GlcNAc residue of N-glycans [11, 13].

FUT8 has been extensively investigated in various cancers. Recent studies indicate that FUT8 expression levels are highly correlated with cancer development, in particular, hepatocellular carcinoma [14], melanoma [15], breast cancer [16], and gastric cancer [17]. Whether FUT8 might also influence the progression of OS is unknown. Furthermore, it is unclear what the underlying mechanism could be.

In the present study, we found significantly lower expression of FUT8 in human OS tissues compared to normal healthy bone tissue. We also found that the expression of FUT8 was negatively correlated with growth of human OS cell lines. In vitro and in vivo assays confirmed that the downregulation of FUT8 in OS cells promoted the progression of OS. Transcriptome sequencing analysis, cell biology experiments and signaling pathway validation provided strong evidence for a possible mechanism. When core fucosylation of TNF receptors was downregulated in OS cell lines due to knockdown of FUT8, the TNF/NF-κB2 signaling pathway was activated and mitochondria-dependent apoptosis was blocked. Together, our findings highlight the important role of FUT8 in the oncogenesis of OS and suggest a new therapeutic strategy for inhibiting OS by changing its glycosylation profile, especially core fucosylation.
pathologically diagnosed with osteosarcoma and underwent primary OS resection. Immediately after excision, the samples were frozen in liquid nitrogen to prevent protein and RNA degradation and then stored at −80 °C. Ethics approval was obtained from the Shanghai Sixth People’s Hospital Ethics Committee (YS-2016-064, 24 February 2016).

Cell lines
The human OS cell lines MNNG/HOS (short for MNNG in the following description), U2OS, and 143B and human osteoblast cell line hFOB1.19 were acquired from American Type Culture Collection (ATCC, USA). OS cell lines MNNG and 143B were cultured in DMEM (Corning, USA) at a 37 °C atmosphere containing 5% CO₂. OS cell line U2OS was cultured in RPMI 1640 (Corning, USA) at a 37 °C atmosphere containing 5% CO₂. Osteoblast cell line hFOB1.19 was cultured in D-MEM/F-12 (Gibco, USA) at a 33.5 °C atmosphere containing 5% CO₂. All culture mediums were supplemented with 10% fetal bovine serum (Gibco, USA).

Stable cell line construction
The lentivirus shuttle plasmids containing full-length FUT8 (for over-expressing FUT8) and sirRNA against FUT8 (for knockdown FUT8) (shFUT8-2 was selected according to efficiency) were co-transfected into HEK293T cells with lentivirus packing vectors, respectively. After 48 h, the supernatant of HEK293T, which containing lentivirus were collected, purified and performed titer determination. Then, MNNG cells were infected by the collected lentivirus at the MOI = 10.0. The positive cells were selected by puromycin at the concentration of 1.0 µg/ml after 72 h after infection. Finally, the stable cells overexpressing (MNNG-F8) and knock down FUT8 (MNNG-siF8) gene were verified by RT-qPCR and WB at both mRNA and protein levels. All the information of the vector and sequences of full-length FUT8 and sirRNA against FUT8 has been provided in the Supplementary Table 1.

Total RNA extraction and Real-time Quantitative PCR
Trizol reagent (Invitrogen, USA) was used to extract total RNA from clinical samples and cell lines. Reverse transcription was achieved by a ReverTra First Strand cDNA Synthesis Kit (Invitrogen, USA). Reverse transcription quantitative PCR (RT-qPCR) assays were carried out on an ABI Prism 7900HT real-time system (Applied Biosystems, USA) by applying specifically primers and the SYBR gene PCR master mix (Invitrogen, USA). The 2−ΔΔCt approach was used to calculate the relative mRNA expression of different genes. All primers are shown in Supplementary Table 2.

Western blot and reagent
RIPA solution (EpiZyme, PRC) containing proteinase inhibitor (Invitrogen, USA) was added to culture cells to gain lysis solution. Then the lysates were centrifuged (13000 g/15 min, 4 °C) and discarded precipitate. Equal amounts of collected total proteins were separated by SDS-PAGE and transferred to a PVDF membrane. Giving the membranes a dip in 5% milk at room temperature (RT) for 1 h. Then, proteins were detected by incubated membranes in primary antibodies solution at 4 °C overnight. After that, HRP-linked anti-IgG antibodies were employed as secondary antibodies. Primary antibodies were shown in Supplementary Table 3.

Real Time Cellular Analysis (RTCA) and colony formation assay
The cell proliferation ability was measured by RTCA (ACEA Biosciences, USA) according to the manufacturer’s instructions. Particularly, first, 100 µl of culture medium was added to wells and incubation at 37 °C in a cell incubator for 1 h. Then the baseline value was measured by the incubated medium. Finally, the cells were seeded into wells, and make sure that there were 2.5 × 10⁴ cells per well. Cell attachment and cell proliferation were continuously recorded for 6 h and 168 h, respectively. Cell index was used to representing cell attachment and cell proliferation [18]. For colony formation ability detection, cells were incubated in a 6-wells plate for 2 weeks at a density of 1 × 10⁴ cells per well. After being fixed with 4% paraformaldehyde (PFA), the cells were stained with crystal violet. The colonies which contain over 50 cells were recorded and counted by image J software.

Subcutaneous tumor model
Four- to five-week-old female nude mice were raised in the Laboratory Animal Research Centre of Shanghai Sixth People’s Hospital. All experiments were approved by the Animal Research Committee of Shanghai Sixth People’s Hospital. For the subcutaneous xenograft tumor model, ten mice were randomly assigned to two groups (N = 5) and anesthetized by intraperitoneal injection of 1% pentobarbital sodium at a dose of 0.01 ml/g. Then, the cells were harvested from cell culture flasks and resuspended in PBS to a final concentration of 5 × 10⁶/ml. After that, 200 µl cell suspension containing 1 × 10⁶ cells were injected into the nude mouse flank [19]. Researchers blinded to group allocation measured the tumors with callipers until the longest diameter of the largest tumor reached 200 mm, and all tumors were extracted via surgery. The volume of tumors was calculated as the length (mm) × width (mm)².

Cell apoptosis analysis
Cells were cultured in a 6-wells plate. After being treated with 0.05 µM staurosporine (STS) (Sigma, USA) for 24 h [20], cells were washed with a cold Phosphate Buffered Saline (PBS) solution followed by digested with trypsin. Digested cells were processed by the Annexin V-PI kit (Beyotime, PRC) according to the manufacturer’s instructions. The result was measured by flow cytometry.

Mitochondrial membrane potential assay
Cells were cultured in a 6-wells plate. After being treated with 0.05 µM STS for 24 h, cells were washed with a cold PBS solution. Cells were processed by mitochondrial membrane potential assay kit (Beyotime, PRC) according to the manufacturer’s instructions. The images were taken by DM6B fluorescence microscope (Leica, BRD).

RNA-seq and analysis
Total RNA was extracted from MNNG-siF8, MNNG-F8, and control cell lines by Trizol reagent (Invitrogen, USA). The RNA-Seq libraries were synthesized by the TruSeq™ RNA Sample Preparation Kit (Illumina, USA) following standard Illumina guidelines. After purification, the quantification and validation of the libraries were performed by Qubit® 2.0 Fluorometer (Life Technologies, USA) and Agilent 2100 bioanalyzer (Agilent Technologies, USA), and finally, the library was sequenced by Illumina NovaSeq 6000 (Illumina, USA). The library construction and sequencing were performed by Sinotech Genomics Co., Ltd (Shanghai, PRC). The selection criteria of differentially expressed genes (DEG) was a less than five percentage false discovery rate (FDR) and changed expression higher than 1.5 or lower than 0.67 folds. All cell lines were tested three times. The raw RNA-seq data was uploaded to NCBI SRA database. The SRA accession number: PRJNA772908.

Cell nucleus isolation
For detection of the nuclear translocation of p52, the nucleus was isolated for western blot analysis. Cells were cultured in a 100 mm dish, and nuclei were extracted with a Nuclei EZ lysis buffer. Then, the nuclei were resuspended and washed in 4 ml of ice-cold Nuclei EZ lysis buffer. Centrifugation was performed again at 500 g for 5 min at 4 °C, and the nuclei pellet was resuspended in 200 µl of ice-cold Nuclei EZ storage buffer and stored at −70 °C. Then, the expression of p52 was detected in both the nucleus and cytoplasm. PCNA and GAPDH were used as internal references.

Immunofluorescence (IF) analysis
Cells were cultured in coverslips which was placed in a 24-wells plate. When confluent up to 30%, cells were fixed with 4% PFA for 20 min at RT. After being washed with PBS, fixed cells were blocked by incubating in QuickBlock™ (Beyotime, PRC) for 1 h. Then, cells were washed by PBS three times and incubated with rabbit anti-NF kappa p100/p52 (1:200, AF6373, Affinity, PRC) overnight at 4 °C. Subsequently, discarded primary antibody and washed by PBS three times, cells were incubated with Alexa Fluor 488-conjugated goat anti-rabbit antibody at RT for 1 h. PI was used to stain nuclei. Images were taken by Confocal Microscope (Leica Microsystems, BRD).

Immunohistochemical (IHC) analysis
The clinical sample tissues were obtained from Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, and the OS tissues were obtained from subcutaneous tumor model. Both clinical tissues and excised OS tissues were embedded with paraffin. Cutting 4µm sections...
from the paraffin-embedded tissue sample and deparaffinized. After antigen retrieval, the samples were blocked with 5% bovine serum albumin (BSA) at 37 °C for 30 min. Then, samples were incubated with primary antibodies overnight at 4 °C. After that, samples were washed by PBS and incubated in HRP-linked anti-IgG at 37 °C for 30 min. Three times PBS washing was performed again and 3,3′-diaminobenzidine (DAB) was added to stain for 10 min. Finally, undergo counterstained and dehydrated, samples could be covered by cover glass and photographed by DM6B (Leica, BRD).

**Immunoprecipitation (IP) and lectin blot**

IP was performed in the MNNG-siFUT8, MNNG-FUT8, and control cell lines using a Pierce communoprecipitation kit (26149, Invitrogen, USA) and anti-TNFR antibody (Affinity, AF0282). Briefly, 20 μg of the TNFR antibody (Affinity, PRC) was incubated with the AminoLink Plus coupling resin for 120 min at room temperature and covalently coupled. During incubation, the protein lysates of cells were extracted by adding 1 ml cold IP lysis/wash buffer to the culture dish (100 x 100 mm). Then, the antibody-coupled AminoLink Plus coupling resin was washed and incubated with 200 μl of the whole cell lysates overnight at 4 °C. Subsequently, the resin was washed again, and the proteins bound to the antibody were eluted by elution buffer. The core fucosylation of the TNF receptor was examined by lectin blot using Aleuria aurantia lectin (AAL) (B-1395, Vectorlabs, USA) from the extracted proteins. The extracted proteins were separated by SDS–PAGE and transferred to a PVDF membrane. The membranes were dipped in 5% milk at room temperature (RT) for 1 h. Then, the proteins were detected by incubating membranes in AAL solution at room temperature for 2 h. Subsequently, HRP-linked avidin was employed as a secondary antibody.

**Statistical analyses**

All data were analyzed by SPSS 25.0 software and presented as the mean ± SD. The two-tailed Student's t-test was performed to calculate the differences between the experimental group and the control group. The Welch's t-test was performed to calculate the differences between the tumor tissues group and non-tumor tissues group of clinical samples. ns means P > 0.05, * means P < 0.05, ** means P < 0.01, *** means P < 0.001.

**RESULTS**

**Reduced expression of core fucosyltransferase FUT8 in human clinical OS samples and OS cell lines**

To examine the role of FUT8 in the progression of OS, we first analyzed the expression of *FUT8* in clinical samples of normal cancellous bone and OS tissue samples by RT-qPCR. *FUT8* mRNA expression was significantly lower in extracts from OS tissue compared to extracts from adjacent normal bone (Fig. 1A). In OS cell lines (MNNG, U2OS, and 143B) and an osteoblast cell line hFOB1.19, we quantified both *FUT8* mRNA and protein levels using qPCR and western blotting. OS cell lines had lower levels of *FUT8* expression compared to that in the osteoblast cell line hFOB1.19 (Fig. 1B, C). We also examined the distribution of FUT8 by immunohistochemical staining OS tissue sections and adjacent non-tumor sections. Consistent with the RT–qPCR results, visual observation of FUT8 immunostaining indicated that it was less dense in OS tissue sections than in nearby normal bone marrow tissue (Fig. 1D). Taken together, these results suggest that downregulation of *FUT8* may be related to tumorigenesis in OS.

**FUT8 negatively regulates the progression of OS in vitro and in vivo**

Since there was significantly reduced expression of *FUT8* in human OS tissue and cell lines, we hypothesized that *FUT8* negatively regulates the progression of OS. To test this idea, we constructed two *FUT8*-regulated stable OS cell lines: MNNG-F8, which overexpresses *FUT8*, and MNNG-siF8, which knockdown *FUT8*. Both of these lines, as well as their control cell lines, were constructed using lentivirus. Successful overexpression and knockdown of FUT8 was verified when we measured its levels of protein (Fig. S1A, 1B) and mRNA (Fig. S1C, 1D). To examine whether altered expression of FUT8 affects cell function in OS cells, we assessed OS cell growth by Real Time Cellular Analysis (RTCA) [18]. Proliferation was measured in MNNG-F8, MNNG-siF8, and control cell lines. As expected, cell proliferation was significantly reduced in cell lines that over-expressed FUT8. By contrast, cell proliferation was enhanced in cell lines that knockdown FUT8 (Fig. 2A–D). The results from colony formation assays followed the same pattern as the RTCA results. Compared with controls, colony-forming assays showed that colony formation was reduced or increased, respectively, depending on whether FUT8 was over- or under-expressed (MNNG-F8 vs. MNNG-siF8, respectively) (Fig. 2E–H).

The role of FUT8 in OS cell proliferation in vivo was also examined in the nude mice subcutaneous implantation tumor model. Tumors were measured every 3 days until the longest diameter of the largest tumor reached 200 mm. FUT8 over-expression inhibited tumor growth, while FUT8 knockdown enhanced tumor growth (Fig. 2I–L).

We were also interested in the possible role of FUT8 in metastasis in OS. Thus, we measured the migration and invasion characteristics of MNNG-F8 and MNNG-siF8 cells using the RTCA assay. A similar pattern of results was obtained for migration and invasion as for the tumor morphological experiment above. Migration and invasion were enhanced in FUT8-knockdown MNNG-siF8 cells. However, migration and invasion tended to be blunted in FUT8-overexpressing MNNG-F8 cells, but the reduction was not significant (Fig. S2). In summary, these results show that the expression of FUT8 is negatively correlated with cell proliferation and colony formation in OS cells.

**Downregulation of FUT8 inhibits apoptosis in OS cells**

We next examined potential mechanism(s) for FUT8 in OS cell growth. During in vitro maintenance of our cell lines, it became apparent that apoptotic rates in MNNG-siF8 were reduced (data not shown). Thus, we measured cell apoptosis with Annexin V/PI staining in MNNG-siF8, MNNG-F8 and their control cell lines treated with or without 0.05 μM STS, and then we identified cell types with flow cytometry. Flow cytometry revealed a significantly enhanced apoptosis phenotype in the MNNG-F8 group (36.53% vs 30.79%) and a significantly depressed apoptosis phenotype in the MNNG-siF8 group (25.25% vs 31.15%), as compared to control cells (Fig. 3A–C). Western blot analysis of apoptosis-related proteins in MNNG-siF8, MNNG-F8 and their control cell lines treated with or without 0.05 μM STS confirmed that the level of FUT8 expression is related to apoptosis. Regardless of the treatment with or without 0.05 μM STS, the levels of cleaved-caspase 3 and cleaved-PARP in MNNG-siF8 cells were significantly reduced while the levels of these two proteins in MNNG-F8 cells were increased (Fig. 3D–F). Together, these results support the hypothesis that FUT8 regulates OS tumor growth through cell apoptosis.

**FUT8-related apoptosis regulation in OS cells occurs through a mitochondria-dependent apoptotic pathway**

In order to examine a possible mechanism of how FUT8 is involved in cell apoptosis, we identified differentially expressed genes (DEGs) via transcriptome sequencing (RNA-seq). This information can help uncover how FUT8 might regulate oncogenesis in OS. RNA-seq analysis revealed a differential gene expression profile between control and engineered FUT8-regulated OS cells (Fig. S3A, 3B). These DEGs are shown in Fig. 4A, B. There were 5,978 DEGs between the control and the FUT8-overexpressing cell line, whereas there were 2,209 DEGs between the control and the FUT8-knockdown cell line.

Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis was carried out among the DEGs. Besides the various types of genes associated with the N-glycan biosynthesis pathway, genes associated with the highly related pathways of TNF signaling
pathway were identified in both MNNG-F8 and MNNG-siF8 cells (Fig. 4C, D). We examined the expression of primary genes related to TNF/NF-κB signaling pathways via RT-qPCR in the cell lines. These results were consistent with the RNA-seq results (Fig. 4A, B). As Figs. 4E, S3C, and S3D show, TRAF1/2, Gadd45b, c-IAP1, IL-8, ICAM-1, PTGS2, MAP3K5, NFKBIA, RIPK2, and NGFR were upregulated while BCL2A1, c-IAP2, Bcl-xl, TNFSF12, and MAP3K14 were downregulated in FUT8-overexpressing cells (MNNG-F8). We observed the opposite expression pattern for these genes in the FUT8-knockdown cells (MNNG-siF8).

The DEGs we identified were highly related to mitochondria-dependent apoptosis. To test whether apoptosis induced by FUT8 overexpression is indeed mitochondria-dependent, we treated the four cell lines with 0.05 μM STS and then stained them for JC-1 in order to detect mitochondrial membrane potential changes (Fig. 5A). JC-1 is cationic dye that is a proxy for mitochondrial transmembrane potentials (ΔΨM) in healthy and apoptotic cells across multiple cell types [21]. The intensity of green fluorescence (JC-1 monomer) was visibly stronger in MNNG-F8 cells, while red fluorescence (JC-1 aggregates) was stronger in MNNG-siF8 cells compared to controls. The lower ratio of red/green fluorescence (Fig. 5B) found in MNNG-F8 cells indicates a lower mitochondrial membrane potential (i.e., impaired mitochondrial function). By contrast, mitochondrial function appeared to be intact in MNNG-siF8 cells (Fig. 5A, B).

Western blot analysis showed that the expression levels of proteins that protected cells from mitochondria-dependent apoptosis (Bcl-2, Bcl-xl, and BCL2A1) were more highly expressed while pro-apoptosis proteins (cleaved-caspase 8 and cleaved-caspase 9) were lower expressed in the MNNG-siF8 cells compared to control cells (Fig. 5C, D). Expression of these proteins in MNNG-F8 cells showed the opposite pattern to that seen in MNNG-F8 cells (Fig. 5C, D). Together, these results support the hypothesis that OS regulated by FUT8 through the mitochondria-dependent apoptotic pathway and can either increase or decrease the cell apoptosis fate of OS cells depending on the expression level of FUT8.

**Downregulation of FUT8 activates the non-canonical TNF/NF-κB signaling pathway**

FUT8 regulates OS cells’ apoptosis via a mitochondria-dependent pathway. KEGG analysis revealed that the non-canonical TNF/NF-κB signaling pathway was altered. We examined in more detail whether this signaling pathways are activated and control mitochondria-dependent apoptosis after FUT8 was remodeled in OS cells. Previous studies showed that members of the TNFR superfamily could activate the NF-κB2 signaling pathway in immune cells [22, 23]. To examine whether the NF-κB2 signaling pathway might also be activated in OS cells, we measured the expression of key proteins involved in NF-κB2 signaling. As shown in Fig. 6A, the knockdown of FUT8 in MNNG-siF8 cells results in the activation of NF-κB2 signaling pathways. Upregulation of NF-κB-inducing kinase (NIK), led to the phosphorylation of p100 (p-p100) and then degraded into p52. By contrast, the overexpression of FUT8 in MNNG-F8 cells inhibited the activation of NF-κB2 signaling.
Since p52 could only play its role as a transcriptional regulator when in nucleus, we next examined the translocation of p52. Western blot analysis detected the expression of p52 both in the cytoplasm and in isolated nuclei. Compared to control cells, there was more p52 detected in the nucleus of FUT8-knockdown cells and less p52 in the nucleus of FUT8-overexpressing cells (Fig. 6B, C). Translocation of p52 into nucleus was also examined by p52 immunofluorescence staining of cultured OS cell lines. The knockdown of FUT8 significantly enhanced the translocation of p52 into nucleus.

Fig. 2  FUT8 expression is negatively correlated with OS oncogenesis in vitro and in vivo. A, C Mean ± SD (N = 4) cell index, a measure of cell attachment and cell proliferation in vitro, for two FUT8-regulated OS cell lines (MNNG-F8, MNNG-siF8) and their control cell lines (F8Control, siControl, respectively) over time. RTCA assay confirmed that cell proliferation is enhanced in vitro in MNNG-siF8 cells, while it is reduced in MNNG-F8 cells compared to control cells. B, D Mean ± SD (N = 4) cell index at 48 h, 72 h, 96 h, 120 h. E, G Representative light microscopic images of PFA-fixed MNNG-F8, MNNG-siF8, and control cells stained with crystal violet, and (F, H) quantitation of colony counts after 2 weeks of in vitro maintenance, data were shown as mean ± SD (N = 3). I, K Images of excised flank tumors after injecting 200 μl of cell suspension from MNNG-F8 (N = 5), MNNG-siF8 (N = 5), or control cells (N = 5) in the nude mouse subcutaneous tumor model, and (J, L) quantitation of tumor volume (mean ± SD). *P < 0.05; **P < 0.01; ***P < 0.001.

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p52 into the nucleus. However, the aggregation of p52 in the nucleus was inhibited by FUT8 overexpression (Fig. 6D, E).

Lack of core fucosylation of TNFR shifts TNF signaling
We were also interested in examining possible changes in signaling upstream of NF-κB. Figure 7A shows a diagram of the method we used for detecting the relative core-fucosylation level of TNFR in F8Control cells, MNNG-F8 cells, siControl cells, and MNNG-siF8 cells. F Quantitative expression values of the WB results in D and E were normalized to β-tubulin expression (N = 3). *P < 0.05; **P < 0.01.

Fig. 3  Expression of FUT8 is positively correlated with the apoptotic rate of OS cells. A, B Two-label dot plots of flow cytometry analysis of two FUT8-regulated OS cell lines (MNNG-F8, MNNG-siF8) and their control cell lines (F8Control, siControl, respectively) treated with either 0 μM or 0.05 μM STS for 24 h. Cells were stained with Annexin V/PI. Compared to control cells, in MNNG-F8 and MNNG-siF8, apoptosis was enhanced and reduced, respectively. C Quantitation of flow cytometry analysis, data were shown as mean ± SD (N = 3). D, E Western blot probed with antibodies against apoptosis-related proteins in the four cell lines with or without 0.05 μM STS treatment confirmed that, compared to control cells, cleaved-caspase 3 and cleaved-PARP were more or less abundantly expressed, respectively, in MNNG-F8 and MNNG-siF8 cells. F Quantitative expression values of the WB results in D and E were normalized to β-tubulin expression (N = 3). *P < 0.05; **P < 0.01.
knockdown cells but was elevated in FUT8-overexpressing cells (Fig. 7B, C). A question is would the core-fucosylation level influence the function of TNFR?

Western blotting was used to examine the expression of Fas-associated death domain (FADD) and TNF receptor-associated death domain (TRADD), which serve as major downstream factors of TNFR to transduce survival and apoptosis signals [24, 25]. A slightly increased and an obviously decreased expression of FADD was detected in the FUT8-overexpressing and FUT8-knockdown cell lines, respectively. However, we found no apparent differences in the expression of TRADD after the fucosylation level of TNFR was changed (Fig. 7D). In summary, these data, along with the RNA-seq results, support our hypothesis that in OS cells, knockdown of FUT8 reduces core fucosylation of TNFR, causing TNFR to transduce signals via FADD to activate TNF/NF-κB signaling pathways.

**FUT8 negatively regulates TNF/NF-κB2 signaling pathway and enhances anti-apoptosis processes in OS**

We demonstrated that FUT8 can alter the core-fucosylation level of TNFR, leading to the activation of the TNF/NF-κB2 signaling pathway. These changes finally affect the phenotype of apoptosis.

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OS tissue from the subcutaneous tumor nude mouse model. IHC results confirmed that, compared to control tissue, MNNG-siF8 tissue showed reduced expression of FUT8, FADD, leading to enhanced expression of p52 and Bfl-1 and reduced expression of cleaved-caspase3. There was an opposite expression trend of these key proteins in MNNG-F8 tissue compared to control tissue (i.e., the enhanced expression of FUT8, FADD, and cleaved-caspase3, and the reduced expression of p52 and Bfl-1) (Fig. 8A).

**DISCUSSION**

Previous studies demonstrated that in OS, a specific glycosylation profile strongly predicts OS progression. For example, Zhang et al., reported that a glycosyltransferase called N-GalNAc transferase is highly expressed in OS [26]. N-GalNAc transferase catalyzes mucin-type O-glycosylation, a common form of glycosylation in mammals [27]. When the N-GalNAc transferase gene is knocked down in OS cells, the invasiveness of OS is significantly reduced...
Deng et al. also showed that loss of O-GlcNAcylation via degradation O-GlcNAc transferase results in the reduction of OS growth [28]. Also, the reduction of galactosyltransferase causes the accumulation of collagen type I in OS, which impaired the OS [29]. However, as far as we know, even though the role of FUT8 has been investigated in many cancers such as lung cancer, prostate cancer, and breast cancer [16, 27, 30–32], the relationship between FUT8 and the progression of OS has received no investigative attention before our study.

In our present study, we demonstrated that the expression of FUT8 was low in OS tissues and OS cell lines compared to that in normal controls. We discovered a negative correlation between FUT8 expression and OS proliferation rate. The knockdown of FUT8 in human OS cell lines resulted in reduced core fucosylation of TNFR, which in turn led to the activation of the NF-κB2 signaling pathway. After being translocated into the nucleus, p52, a NF-κB2 family member, remodeled the expression of mitochondria-dependent apoptosis-related proteins, resulting in inhibition of cell apoptosis (Fig. 8B). The TNF/NF-κB2 signaling pathway is a well conserved, and relatively thoroughly investigated pathway [33]. In addition, the idea that glycosylation plays an important role in protein function has been around for decades [34]. However, most studies on the TNF/NF-κB signaling pathway have focused on the binding between ligands and TNFR and on the phenotypic changes of cells caused by the binding [35–38]. To the best of our knowledge, the present study is the first to link the core fucosylation and the function of TNFR that reducing core fucosylation of TNFR via knockdown of FUT8 leads to activation of the TNF/NF-κB2 signaling pathway.

In line with our main finding, Geert et al. recently presented a novel theory that glycosylation serves as a key regulator of TNFR signaling [39]. Their study showed that N-glycosylation of TNFR was closely related to TNFR's ligand-receptor binding ability, and that cells could change their TNFR's glycosylation profile to

Fig. 6 Knockdown of FUT8 in OS cells in vitro activates TNF/NF-κB2 signaling pathway. A Western blot of proteins in a NF-κB2 signaling pathway show that this pathway was activated in MNNG-siF8 cells but inhibited in MNNG-F8 cells. B Analysis of p52 translocation in OS cell lines. Western blot probed for both in nuclear p52 and cytoplasmic p52. C Quantitation of ratio of nuclear p52 to cytoplasmic p52 expression. Mean ± SD (N = 3) ratio expression in MNNG-siF8 cells was higher, compared to controls, and lower in MNNG-F8 cells. Relative expression of nuclear p52 was normalized to PCNA while relative expression of cytoplasmic p52 was normalized to β-tubulin. D Representative p52 immunofluorescent images of cultured OS cell lines showed translocation of p52 into the nucleus. Red fluorescence represented nuclei, green fluorescence represented p52 (Scale bar: 25 μm). The translocation of p52 into the nuclei appeared visually to be enhanced in MNNG-siF8 cells but reduced in MNNG-F8 cells compared to control cells. E Quantitation of ratio of nuclear p52 to cytoplasmic p52 in cultured OS cell lines. Mean ± SD (N = 3) ratio of nuclear p52 fluorescence to cytoplasmic p52 fluorescence. *P < 0.05; **P < 0.01; ***P < 0.001.
adapt to different microenvironments [39]. This notion of adjusting glycosylation profile according to the microenvironment is consistent with the findings of Holdbrooks et al., who observed that ST6Gal-I sialyltransferase protects tumor cells from TNF-induced apoptosis [40]. They demonstrated that ectopically overexpressed ST6Gal-I sialylated TNFR, which in turn caused the signal induced by TNF to change from an apoptotic one to a survival one [40]. Our results provide further evidence to support this theory that glycosylation, especially core fucosylation, may regulate the function of TNFR. It is also noteworthy that the aberrant expression of FUT8 is commonly found in a variety of tumor cells [13]. Our present results shine a light on the relationship between core fucosylation and membrane receptors in cancers.

About the mechanism, although only a few studies have explored the influence of core fucosylation on membrane receptors in recent years, previously reported findings have important implications for us to solve these problems. For example, normal core fucosylation was demonstrated to be indispensable for the expression of PD-1 on immune cell membranes [41]. PD-1 has long been known to play an important role in how tumors evade the immune system [42]. Likewise, core fucosylation is also required for T-cell receptor transport to their proper place in lipid rafts in T-cell membranes and to transmit T-cell signals properly [43]. In addition, core fucosylation facilitates the binding of TGF-β receptors with their ligands in tumor cells and the phosphorylation of epidermal growth factor receptor in cancer-associated fibroblasts, which could promote the formation of epithelial-mesenchymal transition and tumor microenvironment, respectively [16, 44]. These breakthrough discoveries provide a powerful theoretical foundation for our conclusions and offer hints to some potential mechanisms. In particular, the alteration of FUT8 may play a role in TNFR by affecting their formation and transport, the ability to bind to ligands, and signal transduction, among others.

Although our present findings have illuminated the fairly complete organization and relationships of FUT8, core fucosylation of TNFR, regulation of apoptosis, and the NF-κB signal in OS

Fig. 7 Lack of core fucosylation of TNFR shifts TNF signaling. A Diagram illustrating the method for detecting core-fucosylation level of TNFR in FUT8 regulated cells and their according control cells. TNFR was first enriched from the four cell strains by immunoprecipitation using an anti-TNFR antibody. Then, Western blot and lectin blot, respectively, were used to detect expression of TNFR and core-fucosylation level of TNFR. The lectin blot was done using the AAL, which specifically identifies a combination of core-fucosylated structures. Relative expression of core fucosylation was normalized to TNFR expression. B Western blot of TNFR and lectin blot of core fucosylation of TNFR. Core fucosylation per unit TNFR was increased in MNNG-F8 but reduced in MNNG-siF8 cells compared to control cells. C Quantitation of core-fucosylation levels (N = 3). D Western blot analysis of the shifts TNF signaling. The TNFR down-stream adaptor molecules that mediates cell apoptotic FADD and TRADD were detected by Western blot. The expression of FADD was obvious changed after altering FUT8 levels, while that of TRADD did not change. Relative expression of proteins was normalized to GAPDH. *P < 0.05.
progression. There is still something should be done in the following studies. Firstly, although our hypothesis has been fully verified in the OS cell and pack of patients, it would be more convincing that this signaling pattern could be widely examined in other OS cell lines and more patients. Secondly, the precise mechanisms involved in this loop especially the cascade between the TNFR and the NF-κB2 signaling remain unclear and need to be further studied. Last but not the least, the present research pointed out the important role of the NF-κB2 signaling pathway in the progression of OS. However, until now, no well-accepted therapeutic inhibitor of this pathway has been applied in clinic. Thus, more studies on FUT8 or other potential antagonist of NF-κB2 signaling should be studied to develop the therapeutic inhibitors in the future.

In general, OS remains one of the fatal malignant tumors affecting the quality of life and health of adolescents, thus, there is an urgent need to improve the curative effects of existing therapies [45]. Although the mechanism underlying alteration of the NF-κB2 signaling pathway following core fucosylation of TNFR should be further studied, our present results show that high expression of FUT8 which result in the suppress of NF-κB2 signaling pathway in OS cells is associated with tumor suppression. Conceivably, increasing the expression of FUT8 or suppress the NF-κB2 signaling pathway in tumor cells of patients with OS may provide a new approach to attenuate the tumor burden of OS in the future.

DATA AVAILABILITY
The transcriptomics data generated during this study have been uploaded to NCBI SRA database. The SRA accession number is PRJNA772908. All other data related to this study are available upon request (zhoushumin_zw@126.com).

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AUTHOR CONTRIBUTIONS

Shanyi Lin carried out the experiment and wrote the manuscript; Lenian Zhou carried out the experiment; Yang Dong and Qingcheng Yang prepared the clinical samples; Hanqiang Jin and Quanjun Yang quantified and analyzed the data; Ting Yuan and Shumin Zhou designed the research, reviewed and edited the manuscript. All authors approved the final version of the manuscript. This work was supported by the National Natural Science Foundation of China (NNSFC) (Grant no. 81572239).

COMPETING INTERESTS

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

ADDITIONAL INFORMATION

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