Sclerostin Suppression Facilitates Uveal Melanoma Progression Through Activating Wnt/β-Catenin Signaling Via Binding to Membrane Receptors LRP5/LRP6

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Objective: Uveal melanoma (UM) is the most frequent primary eye cancer in adults with a 50% mortality rate. Characterizing the fundamental signaling pathways that drive UM is of importance for the development of targeted therapy. This study aims to probe the impact of sclerostin (SOST) on malignant progression of UM and regulation of Wnt/β-catenin signaling.

Methods: Epithelial-type (n=20) and spindle-type (n=16) UM tissues were collected for immunohistochemical staining of SOST, Wnt-1, and β-catenin expressions. SOST was silenced in three UM cell lines (primary spindle-type OCM-1 cells, metastatic epithelial Mum-2B cells, and metastatic spindle-type Mum-2C cells) through transfecting specific siRNA. RT-qPCR and Western blot were presented for examining the levels of SOST, and markers in Wnt/β-catenin signaling. Flow cytometry, MTT, EdU, transwell, and tube formation assays were conducted, respectively. By implanting BALB/c nude murine models in situ, the function of SOST on tumor growth was investigated, followed by immunofluorescence double staining of SOST and LRP5/6.

Results: Low SOST expression as well as high Wnt-1 and β-catenin expressions were found in epithelial-type (high malignancy) than spindle-type (low malignancy) UM tissues. Silencing SOST activated the markers in Wnt/β-catenin signaling as well as accelerated cell cycle progression, migration, invasion, angiogenesis, and reduced apoptosis in UM cells. In situ tumor formation in murine eyes showed that SOST knockdown promoted tumor growth. Moreover, SOST interacted with LRP5/6.

Conclusion: SOST silencing may facilitate the malignant progression of UM cells through activating Wnt/β-catenin signaling. Mechanistically, SOST may exert this function by interacting with LRP5/6 membrane receptors.

Keywords: uveal melanoma, sclerostin, Wnt/β-catenin signaling, LRP5/LRP6, progression
INTRODUCTION

Uveal melanoma (UM) represents the most frequent primary intraocular tumor in adults as well as the second most frequent form of melanoma (1). It originates from the pigmented uveal tract, composed of choroid (~85%), ciliary body (5-8%), and iris (3-5%) (2, 3). In accordance with the WHO classification of histological types in 1980, UM is categorized as spindle cell type with low malignancy, epithelioid cell type with high malignancy, mixed cell type, and other types (4). Current therapeutic strategies include surgery, radiotherapy, and chemotherapy as well as targeted therapy, and the local control rate of UM has reached as high as 96.4% following active treatment (5). Nevertheless, approximately 50% of cases will develop metastasis within 15 years following the initial diagnosis (6, 7). Regrettably, there is still no effective treatment against metastasis (8). UM possesses an extremely high predilection to liver metastasis, and exceeding 90% of metastatic patients have hepatic lesions (9). At present, the prognosis of tumor invasion and metastasis cannot be accurately and objectively predicted in clinical practice, and there are not many corresponding effective treatment methods, which are the main factors affecting the curative effect and patient survival rate of UM (10). Therefore, it is necessary to conduct in-depth research on the mechanism of tumorigenesis and tumor metastasis, and to clarify the molecular biological characteristics of this disease.

Wnt/β-catenin signaling exerts a crucial role in tumor progression (11), and blockage of this signaling may suppress growth, migration, and invasion of UM cells (12–14). Sclerostin (SOST) secreted by osteocytes is a negative regulator of bone formation, and is an antagonist of the DAN/Cerberus protein family of BMP (15). Evidence suggests that SOST acts as a potential antagonist of Wnt/β-catenin pathway and inactivated SOST leads to the hyper-activation of this signaling (16). Several studies have reported that SOST-mediated Wnt/β-catenin pathway participates in retinoblastoma progression (17) and osteosarcoma (18). Evidence suggests that SOST can antagonize Wnt signaling in human cells via binding to the extracellular domain of LRPS/6 Wnt co-receptors as well as destroying Wnt-triggered Frizzled-LRP complexes (19). Nevertheless, the roles of SOST-triggered Wnt/β-catenin pathway in UM development as well as its clinical implication remain unclear. Hence, our study was conducted for probing the impact of SOST on malignant progression of UM and regulation of Wnt/β-catenin signaling.

MATERIALS AND METHODS

Patients and Specimens

Twenty epithelial-type UM tissue specimens and 16 spindle-type UM tissue specimens were retrieved from UM patients who underwent enucleation in Tianjin Medical University Eye Hospital from October 2015 to April 2018. All patients were confirmed by pathological examination after operation, who were classified according to the 1980 WHO classification of histological types. The age range was 24-67 years. The inclusion criteria were as follows (1): the pathological diagnosis was primary UM; (2) the submitted tissue sections were relatively complete; (3) the patient had no other types of cancer; and (4) the patient had not received radio- or chemotherapy. This study was approved by the Ethics Committee of Tianjin Medical University Eye Hospital (2017KY (L)-53). All patients provided written informed consent.

Immunohistochemistry

The patients’ eye tumor specimens were fixed in neutral formalin and embedded in paraffin, followed by sectioning, deparaffinization, antigen retrieval, blocking, antibody incubation, DAB (diaminobenzidine) staining, hematoxylin staining, back blue, and dehydration coverslips. Primary antibodies were anti-rabbit (Proteintech, China), and secondary antibodies were goat anti-rabbit IgG (H+L) (Jackson Immunoresearch Inc., US). As previously described, the staining scoring was calculated according to the following formula: H-score = staining intensity * number of positive cells/total number of cells (20). Three random fields were chosen, and the staining intensity was manually evaluated: 0 point for no yellow precipitate, 1 point for light yellow precipitate, 2 points for brownish-yellow precipitate, and 3 points for dark brown-yellow precipitate; positive cell counts were manually counted using Image J software.

Cell Culture

Human retinal epithelial cell line (APRE-19), human umbilical vein endothelial cells (HUVEC) as well as three human UM cell lines were acquired from the Chinese Academy of Sciences (Shanghai, China), including primary spindle-type OCM-1 cells, metastatic epithelial Mum-2B cells, and metastatic spindle-type Mum-2C cells. All cells were cultured in an incubator (Thermo Fisher Scientific, US) with 37°C, 5% CO₂. Subculture was carried out when the cells adhered and reached about 80% confluence. After discarding the medium, they were washed with Dulbecco's Phosphate-Buffered Saline (DPBS; Gibco, US), and digested with 1 mL of 0.25% trypsin-EDTA (Gibco, US). Thereafter, Dulbecco's modified eagle medium (DMEM) containing 100 U/mL penicillin, 100 µg/mL streptomycin, and 10% fetal bovine serum (FBS; Gibco, US) was used to terminate the digestion. The cells were centrifuged at 1000 rpm and resuspended in DMEM for future experiments.

Western Blot

UM cells were lysed by RIPA cell lysate, and protein concentrations were measured utilizing BCA Protein Assay Kit.
GGCGTTCAAGATGATGCCACGGAA-3

5 min. Then, they were resuspended in medium and counted trypsinized at 0.25% as well as centrifuged at 1000 rpm lasting empty vector group and OE-SOST group. These cells were Mum-2C cell lines were randomly divided into two groups: overexpression (OE-SOST) plasmid. OCM-1, Mum-2B, and cloned into the pcDNA3.1 vector to construct pcDNA3.1-SOST plasmid. The full length SOST cDNA was synthesized and sub-cloned into the pcDNA3.1 vector to construct pcDNA3.1-SOST overexpression (OE-SOST) plasmid. OCM-1, Mum-2B, and Mum-2C cell lines were randomly divided into two groups: blank group (no transfection), si-NC group (transfection with si-NC plasmid), and si-SOST group (transfection with si-SOST plasmid). The full length SOST cDNA was synthesized and sub-cloned into the pcDNA3.1 vector to construct pcDNA3.1-SOST overexpression (OE-SOST) plasmid. OCM-1, Mum-2B, and Mum-2C cell lines were randomly divided into two groups: empty vector group and OE-SOST group. These cells were trypsinized at 0.25% as well as centrifuged at 1000 rpm lasting 5 min. Then, they were resuspended in medium and counted using an automated cell counter (Invitrogen, US), and 4×10^5 cells/well were seeded onto a six-well culture plate. When the cell confluence reached 50-80%, the complete medium was replaced with serum- and penicillin-free medium. Four hours later, using Lipofectamine 2000 (Invitrogen, US), plasmids encoding target antisense oligos were transfected into the UM cell line. Twenty-four hours after transfections, the complete medium was exchanged for subculture. During this period, cellular morphology was investigated under a microscope (Olympus, Tokyo, Japan).

**Cell Transfection**

The design and synthesis of the sequences of siRNA targeting human SOST (gene ID: 50964) and negative control siRNA (si-NC) were achieved by Shanghai Sangon Bioengineering Co., Ltd., (China). The sequence of SOST siRNA (si-SOST) was 5’-GGGGTTCAAGATGATGCCACGGAA-3’. OCM-1, Mum-2B, and Mum-2C 2C cell lines were randomly divided into three groups: blank group (no transfection), si-NC group (transfection with si-NC plasmid), and si-SOST group (transfection with si-SOST plasmid). The full length SOST cDNA was synthesized and sub-cloned into the pcDNA3.1 vector to construct pcDNA3.1-SOST overexpression (OE-SOST) plasmid. OCM-1, Mum-2B, and Mum-2C cell lines were randomly divided into two groups: empty vector group and OE-SOST group. These cells were trypsinized at 0.25% as well as centrifuged at 1000 rpm lasting 5 min. Then, they were resuspended in medium and counted using an automated cell counter (Invitrogen, US), and 4×10^5 cells/well were seeded onto a six-well culture plate. When the cell confluence reached 50-80%, the complete medium was replaced with serum- and penicillin-free medium. Four hours later, using Lipofectamine 2000 (Invitrogen, US), plasmids encoding target sequences or siRNAs were transfected into the UM cell line. Twenty-four hours after transfections, the complete medium was exchanged for subculture. During this period, cellular morphology was investigated under a microscope (Olympus, Tokyo, Japan).

**Real-Time Quantitative Polymerase Chain Reaction (RT-qPCR)**

Total RNAs of UM cells were extracted utilizing Trizol kit (Invitrogen, US). The extracted RNAs were utilized for reverse transcription. Cellular cDNA synthesis was conducted according to the instructions of the reverse transcription kit (Transgen, Beijing, China). The PCR amplification reaction was carried out according to the instructions of RT-qPCR kit (Transgen, Beijing, China). Table 1 lists the primer sequences that were synthesized by Shanghai Sangon Bioengineering Co., Ltd. (China). The reaction condition contained pre-denaturation at 94°C lasting 4 min, denaturation at 94°C lasting 30 sec, annealing at 56°C lasting 30 sec, and extending at 72°C lasting 30 sec, in total 45 cycles. The reaction system contained 10 μl Premix Ex Taq or SYBR Green Mix, 1 μl forward primer, 1 μl reverse primer, 1 μl cDNA template, and ddH₂O supplemented to 20 μl. Glyceraldehyde phosphate dehydrogenase (GAPDH) acted as an internal reference gene, and the Ct values (threshold of amplification curves) were examined with the control group set to 1. The iQ5 Real-Time PCR Amplifier was purchased from the Bio-Rad company (US). Relative expression values were calculated by the 2^ΔΔCt method.

**Cell Cycle Detection by Flow Cytometry**

UM cells were collected and digested with trypsin, followed by centrifugation at 1000 rpm for 5 min. After rinsing the cells with sterile PBS, 1 ml of pre-cooled 70% ethanol was utilized to resuspend the cells as well as fixed overnight at 4°C. They were then rinsed with PBS to remove the ethanol solution, followed by incubation with 500 μl of PI staining solution at room temperature lasting 15 min in the dark. The cell cycle was detected by flow cytometry.

**Aptoptosis Detection via Flow Cytometry**

UM cells were trypsinized, followed by centrifugation at 1000 rpm lasting 5 min. After rinsing the cells utilizing pre-cooled sterile PBS at 4°C, the cellular concentrations were adjusted to 5×10⁶/mL. Then, 10 μl of Annexin V-FITC and 10 μl of PI were added to 200 μl of the cell suspension and incubated for 10 min at room temperature in the dark, followed by the addition of 500 μl PBS. Apoptotic levels were examined by flow cytometry (BD, US) as well as analyzed with FlowJo flow cytometry software.

**Methyl Thiazolyl Tetrazolium (MTT)**

Following adjustment of the cellular concentration to 1×10⁷/mL, the cells were seeded in 96-well plates, with six duplicate wells for each group. The edge wells were filled with 0.01 mol/L sterile PBS. After 24 h at constant temperature, the medium in the wells was aspirated and discarded, and 90 μl sterile PBS supplemented with 10 μl 0.5% MTT (Sigma-Aldrich, US) were added. After 4 h, the culture was terminated. Thereafter, 100 μl of dimethyl sulfoxide (DMSO; Sigma-Aldrich, US) was added to each well. The plate was shaken well for 10 min, and the absorbance (optical density, OD) value at 490 nm wavelength of each well was measured with a microplate reader (Promega, US).

| Gene name | Primer sequence (5′-3′) |
|-----------|------------------------|
| SOST      | Forward: TGAGACAAAGACGAGTTCGCG  |
| WNT1      | Reverse: CTTTGAGCTCACTGGTTG  |
| CTNB1     | Forward: CGTGAGTGTTGGATTTGAC  |
| CCND1     | Reverse: CGCGTATTGGGATGATGAC  |
| MMP2      | Forward: GCTGCGGAATGGACGACACTC |
| MMP9      | Reverse: CTCTCTGTCACACATTGGA  |
| GAPDH     | Forward: TACAGGCTATGTTGACAACC |
|           | Reverse: GGCGGAGATTCCTCCAAAAT |

The primer sequence utilized for RT-qPCR.

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**Table 1** The primer sequence utilized for RT-qPCR.
**EdU Staining**

UM cells were seeded in 24-well plates (1 mL/well) containing 1 cm × 1 cm sterile glass slides. The plates were pre-incubated for 24 h at 37°C in a 5% CO2 incubator. After transfection, 20 μM EdU working solution pre-warmed at 37°C was added to a 24-well plate. The cells continued to incubate for 2 h. After removing the culture medium, the cells were fixed with 1 ml of 4% paraformaldehyde for 15 min at room temperature. After washing, the cells were incubated with 1 ml of 0.3% TritonX-100 in PBS per well lasting 10 min at room temperature, followed by 0.5 ml Click additive reaction lasting 30 min in the dark. DAPI (1000×) was diluted 1:1000 in PBS. After removing the washing solution, they were incubated with 200 μl of 1× DAPI solution at room temperature lasting 5 min in the dark. Finally, fluorescence detection was performed.

**Transwell Assays**

For invasion assay, Matrigel was diluted 8:1 with serum-free DMEM. Then 100 μl diluted Matrigel was added to the upper chamber of the transwell, and the whole process was performed on ice to prevent it from solidifying. The residual liquid was aspirated from the culture plate. UM cells were trypsinized, resuspended, and concentration adjusted. The upper chamber of a 24-well plate transwell covered with Matrigel was evenly seeded 1×10^5 cells/well as well as maintained by adding serum-free DMEM. Meanwhile, 600 μl of complete medium with 10% FBS was added to the lower chamber. Following 24 h, the chamber was removed, rinsed twice with PBS, and then gently wiped with a cotton swab to remove unpenetrated cells on the upper layer of the chamber membrane. The cells were fixed with 4% paraformaldehyde for 30 min in a 24-well plate, stained with crystal violet staining solution lasting 15 min. The cells invaded through Matrigel were observed under an inverted microscope. Three areas were randomly selected to count the number of invasive cells through Matrigel. The experimental procedure for the migration assay was the same as for the invasion assay, except that Matrigel was not added.

**Tube Formation Assay**

UM cells transfected with si-NC or si-SOST were co-cultured with HUVEC, and the tube formation ability was detected. Briefly, Matrigel was removed from -20°C and placed in a 4°C refrigerator overnight. The next day, 50 μl Matrigel was added to pre-cooled 96-well plates, and maintained in a 37°C incubator for 45 min. The co-cultured HUVEC was routinely digested, resuspended in cell medium containing 10% FBS. Thereafter, they were seeded into 96-well plates (1×10^4 cells/well) as well as incubated at 37°C. The tube formation was investigated under an inverted microscope.

**Animal Experiment**

The primer of SOST was designed by querying the CDS sequence on the NCBI website as well as the Thermo website, with a primer sequence of 5'- AAAAGCAGGGTTCAAGAATG ATGCTTGGATGCCAAGCATTTGAAACGCCTGC-3'. The primer was synthesized by Shanghai Sangon Bioengineering Co., Ltd. (China). With pENTRM H1/TO as vector, shRNA SOST (sh-SOST) or negative control (sh-NC) plasmids were constructed by enzyme digestion, ligation, transformation, and other operations, and then the plasmid identification was carried out. There were 293 T cells used to package the target plasmid, and when the density of OCM-1 cells reached 40%-50%, a stable transfection cell line was constructed after screening.

Six-week-old BALB/c nude female mice (14 ± 2 g) were purchased from Vital River Company (Beijing, China). All experiments gained the approval of the Laboratory Animal Care and Use Committee of Tianjin Medical University Eye Hospital (TJJY2020122054). All animals were separated into four groups (3 mice/group), containing a control group, model group, model + sh-NC group, and model + sh-SOST group. Normal OCM-1 cells or OCM-1 cells stably transfected with sh-NC or sh-SOST were trypsinized, centrifuged as well as suspended in Hanks’ balanced salt at 1×10^7/mL. The abdominal skin of the mice was disinfected with iodophor, and the mice were anesthetized by intraperitoneal injection of 0.5% pentobarbital sodium. The right eye of the mice was instilled with proparacaine for ocular surface anesthesia, and the left eye was coated with ophthalmic gel to prevent keratitis. Under the operating microscope, a 30G needle was pointed at the oblique angle to create a tunnel via the sclera, and to penetrate the choroid without retina. There were 34G blunt needles attached to 10 μl Hamilton syringes inserted into the subretinal space with the bevel toward the inner eye. The mice in the model group, model + sh-NC group, and model + sh-SOST group were separately injected by 5 μl (5×10^4 cells) of the cell suspension of normal OCM-1 cells or OCM-1 cells stably transfected by sh-NC or sh-SOST. For the control group, the mice did not receive any treatment. After the injection, a white bump was found in the fundus of the mouse, indicating that the injection was successful. Tobramycin ophthalmic ointment was then applied to the eye to reduce the risk of infection. After successful injection, the state of the mice was observed and weighed every other week, and the time of tumor formation in the eyes of the mice was recorded. After six weeks, all mice were euthanized, and their eyeballs were enucleated. The length (L) as well as width (W) of the murine eyeballs were measured utilizing a vernier caliper. The tumor volume (V) was measured in accordance with the following formula: $V = \pi \times L \times W^2 \times \frac{1}{6}$.

**TUNEL Staining**

A TUNEL kit was purchased from the Roche Company (Switzerland). The tissue slides were soaked in xylene for 10 min to dewax. For hydration, the sections were soak in graded ethanol (100%, 95%, 85%, 70%, 50%) for 30 sec, respectively. After washing twice with PBS for 5 min each, 20 μg/ml proteinase K was dropwise added. After 15 min at 37°C, the sections were washed 3 times with 1×PBS. The labeling solution was mixed with buffer evenly at a volume ratio of 1:9. Thereafter, 100 μL TUNEL reaction solution was dropwise added. The reaction area with a cover glass was covered, and reacted at 37°C in a humid, dark environment for 60 min. After washing three times with 1×PBS, 50 μL of DAPI buffer (0.1 mg/mL; Sigma, US) was added to each well, and incubated in the dark at room temperature for 5 min. After washing 3 times with 1×PBS, images were photographed under a fluorescence microscope.
**Immunofluorescence**

Paraffin sections of murine *in situ* UM tumors in eyeball were deparaffinized, antigen retrieved, blocked with 5% BSA, incubated with primary and secondary antibodies, and finally stained with DAPI (Sigma, US) and mounted with resin. Primary antibodies included SOST, LRP5, and LRP6 antibodies (Affinipure, Jackson, US), and fluorescent-labeled secondary antibodies included Alex Fluor 488 and Alex Fluor 610 goat anti-rabbit antibodies (Affinipure, Jackson, US). The sections were then observed under a fluorescence microscope and three fields of view were randomly selected to be photographed. The nuclei stained with DAPI were blue under the excitation of ultraviolet, and the positive expression was fluorescein-labeled green or red.

**Statistical Analysis**

All statistical analysis was carried out utilizing GraphPad Prism software (version 8.0.1, GraphPad Software, US). Student’s t-test was implemented for comparisons between two groups, and one-way analysis of variance (ANOVA) was implemented for comparisons between multiple groups. Pearson correlation test was used for correlation analysis. P<0.05 was considered statistically significant.

**RESULTS**

**Down-Regulated SOST and Up-Regulated Wnt-1 and β-Catenin in Epithelial Type Than Spindle Type UM Tissues**

Our study analyzed the expressions of SOST and Wnt-1 as well as β-catenin in 20 cases of epithelial type UM tissues with a high degree of malignancy and 16 cases of spindle type UM with a low degree of malignancy via immunohistochemistry. The results showed that SOST expression was lower as well as Wnt1 and β-catenin expressions were higher in epithelial type in comparison to spindle type UM tissues (*Figures 1A, B*). This indicated that low SOST expression as well as activation of Wnt/β-catenin signaling were linked to malignant progression of UM. We also calculated the correlation between SOST and Wnt1 and β-catenin in UM. Our results showed that SOST was negatively correlated to Wnt1 (p=0.0003 and r=-0.9869) and β-catenin.
SOST Knockdown Activates Wnt/β-Catenin Signaling in UM Cells

SOST expression was further examined in human retinal epithelial cell line (APRE-19) and three human UM cell lines containing primary spindle-type OCM-1 cells, metastatic epithelial Mum-2B cells, and metastatic spindle-type Mum-2C cells. As shown in Figures 2A, B, SOST expression was lower in three human UM cell lines in comparison to APRE-19 cells. Compared with OCM-1 cells, lower SOST expression was found in Mum-2B and Mum-2C cells. Meanwhile, SOST expression was lower in Mum-2C cells than Mum-2B cells. This indicated the down-regulation of SOST in metastatic cells. To silence the expression of SOST, siRNA against SOST was transfected into OCM-1, Mum-2B, and Mum-2C cells. In Figure 2C, 48 h after transfection, the expression of SOST mRNA was successfully decreased in three human UM cell lines. Further analysis showed that the expressions of WNT1, CTNNB1, CCND1, MMP2, and MMP9 mRNAs were decreased in si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells than controls (Figures 2D–H). As expected, SOST protein was reduced following si-SOST transfection than control in three human UM cell lines (Figures 2I, J). Additionally, Wnt-1, β-catenin, Cyclin-D1, MMP2, and MMP9 proteins presented higher levels when SOST expression was knocked out (Figures 2K–O). Collectively, silenced SOST could activate the Wnt/β-catenin pathway in UM cells.

Low SOST Expression Accelerates Cell Cycle Progression As Well As Reduces Apoptosis in UM Cells

Flow cytometry showed that after si-SOST transfection, the proportion of G1 phase was lower as well as the proportions of S phase and G2 phase were higher in three cell lines OCM-1, Mum-2B, and Mum-2C compared with that of the si-NC group, indicating that silencing SOST expression was capable of increasing the proportion of UM cells in interphase (Figures 3A–F). Additionally, the apoptotic levels of si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells were decreased in comparison to that of the si-NC group (Figures 3G–L). Altogether, SOST knockdown
FIGURE 3 | Low SOST expression accelerates cell cycle progression and reduces apoptosis in UM cells. (A–F) Cell cycle distribution of normal, si-NC-, or si-SOST-transfected OCM-1 and Mum-2B as well as Mum-2C cells. (G–L) Apoptotic levels of normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. *p<0.05; ****p<0.0001.
enabled us to accelerate cell cycle progression as well as decrease apoptosis in UM cells.

**SOST Knockdown Strengthens Proliferative Capacities of UM Cells**

Through CCK-8 as well as EdU staining assays, this research evaluated the effects of SOST on proliferation of UM cells. As depicted in Figures 4A–C, the cell viability of si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells was elevated in comparison to that of the si-NC group. To overexpress SOST, UM cells were transfected with OE-SOST plasmid. RT-qPCR confirmed that SOST expression was significantly up-regulated in UM cells (Figure 4D). Lower cell viability was found in SOST-overexpressed OCM-1, Mum-2B, and Mum-2C cells (Figure 4E). Moreover, proliferation levels of three human UM cell lines were enhanced by si-SOST transfection (Figures 4F, G) as well as weakened by SOST overexpression (Figures 4H, I). Thus, reduced expression of SOST enabled us to strengthen proliferative capacities of UM cells.

**Silencing SOST Facilitates Migration, Invasion, and Angiogenesis of UM Cells**

Transwell experiment results showed that compared with the si-NC group, the number of OCM-1, Mum-2B, and Mum-2C cells that penetrated the upper chamber of transwell in the si-SOST group was elevated in the si-SOST group (Figures 5A, B). This indicated the increase in migration capacities of UM cells. In Figures 5C, D, the number of three human UM cell lines invaded through Matrigel was increased in the si-SOST group in comparison to the si-NC group, indicating the enhancement in invasion capacities of UM cells. Following co-culture with si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells, the tube formation number of HUVEC was increased (Figures 5E, F). Altogether, SOST knockdown could enhance migration and invasion as well as angiogenesis of UM cells.

**SOST Suppression Strengthens Tumor Growth in Murine Models**

To further verify the impact of SOST on UM, we constructed a stable SOST knockdown expression OCM-1 cells as well as ocular orthotopic tumor models in nude mice. Following six weeks, we removed the eyeballs and measured the tumor volume. In comparison to the control group, the tumor volume of the model group was increased. Meanwhile, higher tumor volume was found in the model + sh-SOST group than the model + sh-NC group (Figures 6A, B), indicating that SOST suppression accelerated tumor growth. TUNEL staining also demonstrated that proliferative capacity of tumor cells in the model + sh-SOST group was heightened in comparison to that of the model + sh-NC group (Figure 6C).

**SOST Suppression Activates Wnt/β-Catenin Signaling in Ocular Orthotopic Tumor Models**

Immunohistochemistry demonstrated that SOST expression was lower in tumor cells of the model + sh-SOST group in comparison to that of the model + sh-NC group (Figure 7). Additionally, Wnt-1, β-catenin, cyclin-D1, MMP2, and MMP9 as well as Ki-67 levels were higher in tumor cells of the model + sh-SOST group than that of the model + sh-NC group, demonstrating that SOST suppression was capable of activating Wnt/β-catenin signaling in ocular orthotopic tumor models.

**Interaction Between SOST and LRP5/6 in UM Cells**

Further analysis showed that LRP5 and LRP6 membrane receptors presented lower expressions in tumor cells of the model + sh-SOST group than that of the model + sh-NC group, demonstrating that SOST suppression could decrease the expressions of LRP5 and LRP6 (Figures 8A, B). Also, SOST co-localized with LRP5/6 in UM cells, indicating that SOST was interacted with LRP5/6.

**DISCUSSION**

UM is an aggressive malignancy that originates from ocular melanocytes, which is an incurable melanoma and the mortality rate is as high as 50% (21, 22). The experimental results from this study demonstrated that SOST silencing could heighten the proliferation, migration, and invasion as well as angiogenesis of human UM cells via activating Wnt/β-catenin signaling via binding to membrane receptors LRP5/LRP6.

Low SOST expression as well as high Wnt-1 and β-catenin expressions were found in epithelial-type (high malignancy; n=20) in comparison to spindle-type (low malignancy; n=16) UM tissues. Silencing SOST activated Wnt/β-catenin signaling in three human UM cell lines. Moreover, its knockdown accelerated cell cycle reduced apoptosis as well as facilitated migration, invasion, and angiogenesis of UM cells. SOST is primarily expressed in osteocytes, as a negative regulator of bone formation (23, 24). SOST deficiency facilitates bone formation via Wnt/β-catenin signaling pathway as well as compensates particle-triggered osteolysis (25). Accumulated evidence has demonstrated that SOST mediates cancer progression. Hudson et al. found that SOST participates in the early onset of bone metastasis of prostate cancer, which can inhibit the invasion of prostate cancer cells as well as subsequent bone metastasis (26). SOST deficiency triggers proliferative and invasive capacities and decreases apoptotic levels of human retinoblastoma cells (17) and osteosarcoma cells (18) via activation of Wnt/β-catenin signaling. Hesse et al. reported that suppressing SOST is capable of alleviating breast cancer-triggered bone metastasis and muscle weakness (27). In postmenopausal Romanian women with osteoporosis, there is a close association between LRP5 and SOST polymorphisms (28). Evidence suggests that LRP5/6 co-deletion in bone markedly reduces the skeletal gain from SOST antibody treatment (29). Herein, our immunofluorescence double staining demonstrated the interactions between SOST and LRP5/6 in UM cells, proving that SOST directly bound to LRP5/6 membrane receptors, thereby inhibiting Wnt/β-catenin signaling during UM progression.

To confirm the above demonstration, this study also used the human UM cell line OCM-1 to construct a stable SOST knockdown expression cell line. Thereafter, we established an
FIGURE 4 | SOST knockdown strengthens proliferative capacities of UM cells. (A–C) CCK-8 for the cell viability of normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. (D) RT-qPCR for the expression of SOST in empty vector- or OE-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. (E) CCK-8 for the cell viability of empty vector- or OE-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. (F, G) EdU staining for the proliferation of normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. (H, I) EdU staining for the proliferation of empty vector- or OE-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. Magnification, ×200. Scale bar, 10 μm. **p<0.01; ***p<0.001; ****p<0.0001.
FIGURE 5 | Silencing SOST facilitates migration and invasion as well as angiogenesis of UM cells. (A, B) Transwell experiments for the migration of normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. Magnification, ×40. Scale bar, 10 μm. (C, D) Transwell experiments for the invasion of normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. Magnification, ×40. Scale bar, 10 μm. (E, F) Tube formation experiment of HUVEC co-cultured with normal, si-NC-, or si-SOST-transfected OCM-1, Mum-2B, and Mum-2C cells. Magnification, ×40. Scale bar, 200 μm. **p<0.01; ***p<0.001; ****p<0.0001.
animal model of ocular orthotopic tumor implantation in BALB/c nude mice to verify the effects of SOST inhibition on UM tumor formation. The results showed that when SOST expressions were knocked down in human UM cells, the tumor volume of ocular tumors of nude mice was larger. Additionally, the expressions of Wnt-1, β-catenin, Cyclin-D1, and MMP2 as well as MMP9 were all increased when the expression of SOST was suppressed in UM cells. As demonstrated by Kim et al. SOST is capable of inactivating Wnt signaling in Xenopus embryos (30).

Furthermore, Zhou et al. further proved that the expression of cyclin-D1 is linked with the progression of UM, and high Let-7b expression may weaken the expression of cyclin-D1 and heighten cell cycle arrest to increase the radiosensitivity of UM cells (31). Chang et al. verified that epigallocatechin gallate may downregulate the activity of secreted MMP-2 via suppression of ERK1/2 phosphorylation, and thus reduces the migration of UM cells (33). Cheng et al. demonstrated that HMGA1 can accelerate the progression of UM via positively modulating the activity of PI3K/Akt/MMP9 signaling in xenograft murine models (34). The above evidence demonstrated that SOST suppression may heighten UM progression through activating Wnt/β-catenin signaling via binding to membrane receptors LRP5/LRP6.
CONCLUSION

Taken together, this research proposed that SOST suppression may activate the Wnt/β-catenin pathway, thereby accelerating UM malignant progression. We also have reasons to believe that the SOST-triggered Wnt/β-catenin pathway might represent a promising therapeutic as well as adjunctive regimen against UM. Despite this, future clinical trials are needed for determining the safety as well as effects of this therapy. Due to the small number of subjects, it is difficult to generalize the results. To gain a deeper understanding of the biological behaviors of UM cells as well as better assess the prognosis of patients, clinical trials and clinical studies on a larger population based on the SOST-triggered Wnt/β-catenin pathway are also needed.

FIGURE 7 | SOST suppression activates Wnt/β-catenin pathway in ocular orthotopic tumor models. Immunohistochemistry staining for SOST, Wnt-1, β-catenin, cyclin-D1, MMP2, and MMP9 as well as Ki-67 in tumor sections from mice injected with normal, sh-NC-, or sh-SOST-transfected OCM-1 cells after six weeks. Magnification, ×40. Scale bar, 200 μm.
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material. Further inquiries can be directed to the corresponding authors.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Tianjin Medical University Eye Hospital (2017KY (L)-53). The patients/participants provided their written informed consent to participate in this study. The animal study was reviewed and approved by Tianjin Medical University Eye Hospital (TJYY2020122054).

AUTHOR CONTRIBUTIONS

TW and XH conceived and designed the study. HW and SZ conducted most of the experiments and data analysis and wrote the manuscript. YL and FS participated in collecting data and helped to draft the manuscript. All authors reviewed and approved the manuscript.

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