Long non-coding RNA GHET1 promotes osteosarcoma development and progression via Wnt/β-catenin signaling pathway

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Abstract. Osteosarcoma (OS) is known as a malignant tumor with a high mortality rate of children and adults worldwide. Long non-coding RNAs (lncRNAs) have been revealed as oncogenes or tumor suppressors that are involved in the tumorigenesis and metastasis of some types of cancer. However, the biological role of long non-coding RNA gastric carcinoma proliferation enhancing transcript 1 (lncGHET1) and its regulatory mechanism in OS progression have not been elucidated. The aim of the present study was to investigate the role of lncGHET1 in OS. The present study explored lncGHET1 expression using a reverse transcription-quantitative (RT-q) PCR assay. Furthermore, the Cell Counting Kit-8 assay, flow cytometry detection, wound healing and transwell invasion assays were performed to evaluate its biological role and the underlying mechanisms in vitro. Additionally, the effect of lncGHET1 was evaluated in vivo in a xenograft model. lncGHET1 expression was significantly upregulated in OS cell lines compared with in an osteoblastic cell line according to the RT-qPCR assay. The results of a knockdown functional experiment suggested that inhibition of lncGHET1 attenuated cell proliferation, migration, invasion and epithelial-to-mesenchymal transition, and promoted apoptosis, partly through regulating the Wnt/β-catenin signaling pathway in OS. These findings indicated that lncGHET1 may serve an essential regulatory role in the biological processes of OS. The present study identified a novel therapeutic target for diagnosis and treatment of human OS.

Introduction

Osteosarcoma (OS) is a primary malignant bone tumor that typically affects the long tubular bones of children and adults worldwide (1). OS is characterized by the direct formation of immature bone or osteoid tissue by tumor cells (2), as well as poor prognosis (3,4). Despite medical advances and development of early diagnosis, the prognosis of patients with metastatic osteosarcoma remains poor, with a cure rate of ~30% (5-7). OS remains a leading cause of cancer-associated mortality among children and teenagers (8-10). Therefore, it is urgently necessary to develop novel targets and research the underlying mechanisms of bone carcinogenesis, which is critical for the treatment and prognosis of OS in the future.

Long non-coding RNAs (lncRNAs) are sequences >200 nt in length and without protein-coding capacity (11). Previous studies have provided evidence that lncRNAs serve significant roles in the biological behavior of tumors, including proliferation, invasion, metastasis, differentiation and apoptosis (12-15). Notably, a previous study has confirmed the dysregulation of lncRNAs in human OS (16). lncRNA FGFR3-ASI promotes OS growth by regulating fibroblast growth factor receptor 3 (17). In addition, lncRNA small nucleolar RNA host gene 12 promotes OS cell proliferation and migration in vitro (18). Therefore, further investigations are required to evaluate the expression pattern, biological role and functional mechanisms of lncRNAs in OS. Long non-coding RNA gastric carcinoma proliferation enhancing transcript 1 (lncGHET1; AK123072) has been demonstrated to be abnormally expressed in gastric, bladder and colorectal cancer (19-22). However, the role and underlying mechanisms of lncGHET1 in OS remain unclear.

The present study analyzed the lncGHET1 expression pattern in OS cell lines using reverse transcription-quantitative PCR (RT-qPCR). In addition, loss-of-function experiments were performed to investigate the biological roles of lncGHET1 in OS. The results revealed that lncGHET1 expression was upregulated in OS cell lines. Additionally, the results of in vitro and in vivo assays revealed that silenced lncGHET1 inhibited cell proliferation, tumor growth, migration, invasion and epithelial-to-mesenchymal transition (EMT), and promoted apoptosis, partly via the Wnt/β-catenin signaling pathway. The findings suggested that lncGHET1 functions as an oncogene, which may contribute to the development for diagnosis and treatment of OS.

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Materials and methods

Cell culture and transfection. Human OS U2OS, MG-63 and SaOs-2 cell lines, and the epithelial hFOB cell line were purchased from American Type Culture Collection. All cells were cultured in RPMI-1640 medium (Invitrogen; Thermo Fisher Scientific, Inc.) supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) and 100 U/ml penicillin and streptomycin (Thermo Fisher Scientific, Inc.) and maintained at 37˚C with 5% CO₂. Small interfering RNA (si/siRNA)-IncGHET1 (5'-CGGCGAGCGATGGATACAGA-3') and negative control (si-NC) (5'-CGGCAGAGCCAUGAGCA-3') were designed and synthesized by Shanghai GenePharma Co., Ltd. For cell transfection, 50 nM siRNA or si-NC was transfected into U2OS and MG-63 cells (1x10⁶) using Lipofectamine® 2000 (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer's protocol. After 48 h, the cells were collected for further experimentation. The transfection efficiency was evaluated by GFP and RT-qPCR analyses.

Cell Counting Kit-8 (CCK-8) assay. A CCK-8 (Sigma-Aldrich; Merck KGaA) assay was performed to examine cell viability according to the manufacturer's protocol. Briefly, MG-63 and U2OS cells were seeded at a density of 1x10⁴ per well and incubated in a humidified incubator at 37˚C for 24, 48 and 72 h. Subsequently, the cells were incubated with 10 µl CCK-8 solution for another 2 h at 37˚C. Optical density was determined at a wavelength of 450 nm.

Colony formation and 5-ethyl-20-deoxyuridine (EdU) assay. For the colony formation assay, transfected cells (1,000 cells/well) were seeded and cultured for 14 days. Subsequently, the cells were fixed with 4% polyoxymethylene for 20 min at room temperature and stained with 10% crystal violet for 30 min at room temperature. The colonies were counted and analyzed under a light microscope. For the EdU assay, 1x10⁶ transfected cells were seeded and cultured for 24 h, and then fixed with 4% paraformaldehyde at room temperature for 15 min, followed by 100 µl 1X Hoechst 33342 (5 µg/ml) staining for 30 min at room temperature. Cells were visualized under a fluorescence microscope (Leica Microsystems GmbH).

RT-qPCR analysis. Total RNAs from U2OS and MG-63 cells with different treatments were extracted using TRIzol® reagent (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer's protocol. Reverse transcription was performed with SuperScriptTM II reverse transcriptase (Invitrogen; Thermo Fisher Scientific, Inc.). The protocol was as follows: 42˚C for 60 min, 70˚C for 15 min and chilling at 4˚C. The relative levels of genes were detected by RT-qPCR using SYBR Premix Ex Taq™ (Takara Bio, Inc.). PCR cycling conditions were as follows: 95˚C for 5 min, followed by denaturation for 10 sec at 95˚C and extension for 30 sec at 60˚C for 40 cycles, and 60˚C for 5 min. GAPDH was used as an internal loading control. All reactions were performed in triplicate. Fold changes were calculated using the 2−ΔΔCT method (23). The primers were as follows: GHET1 forward, 5'-CCCCCACAATGAGAGACACT-3' and reverse, 5'-TTC CCAACACCTATAAGAT-3'; snail forward, 5'-TGTTGCA GTGAGGGCAAGAA-3' and reverse, 5'-GACCTCTGGTGCTTCAAGGA-3'; N-cadherin forward, 5'-CGAGCGGCCGC TCGCCTGCCAC-3' and reverse, 5'-CGCTGCTCTCCGGTCC CCCGC-3'; E-cadherin forward, 5'-TACGCCCTGGGACTCT ACCTA-3' and reverse, 5'-CCAGAAACCGAGGCTCTGAT-3'; vimentin forward, 5'-TACAGGAAACGTCTGGAAAG-3' and reverse, 5'-ACCAGGGAGTGAATCCACG-3'; and GAPDH forward, 5'-CGGAAGTCACGGATTGTTGCTAT-3' and reverse, 5'-AGCCTTTCTCATGTTGGTGAAGAC-3'.

Western blot analysis. Total proteins were extracted using RIPA buffer (Cell Signaling Technology, Inc.) and the protein concentrations were determined using the BCA Protein Assay kit (Beyotime Institute of Biotechnology). Proteins (30 µg/lane) were separated by 10% SDS-PAGE and transferred to PVDF membranes (EMD Millipore). Membranes were blocked with 5% non-fat milk for 2 h at room temperature. Subsequently, the membranes were incubated with primary antibody at 4˚C overnight. After washing with TBS with 1% Tween-20, the membranes were probed with HRP-conjugated rabbit anti-mouse IgG secondary antibodies (cat. nos. ab6721 and ab6728; dilution, 1:500; Abcam) for 1 h at room temperature. GAPDH was used as a loading control. The protein bands were visualized using an Enhanced Chemiluminescence Detection system (Thermo Fisher Scientific, Inc.). The data were analyzed using Image-Pro plus software v6.0 (National Institutes of Health). Antibodies for p21 (cat. no. 64016; dilution, 1:1,000), p27 (cat. no. 3688; dilution, 1:2,000), cyclin D1 (cat. no. 55506; dilution, 1:2,000), GAPDH (cat. no. 5174; dilution, 1:5,000), Bax (cat. no. 2772; dilution, 1:2,000), Bcl-2 (cat. no. 3498; dilution, 1:2,000), caspase-3 (cat. no. 9664; dilution, 1:1,000), caspase-9 (cat. no. 9509; dilution, 1:2,000) and MMP-2 (cat. no. 87809; dilution, 1:1,000) were purchased from Cell Signaling Technology, Inc. Antibodies for MMP-9 (cat. no. ab38898; dilution, 1:2,000), Snail (cat. no. ab53519; dilution, 1:2,000), N-cadherin (cat. no. ab18203; dilution, 1:2,000), E-cadherin (cat. no. ab231303; dilution, 1:2,000), vimentin (cat. no. ab92547; dilution, 1:1,000), Wnt (cat. no. ab219412; dilution, 1:1,000) and β-catenin (cat. no. ab6051; dilution, 1:1,000). All of these antibodies were purchased from Abcam.

Wound healing assay. In order to evaluate the role of lncGHET1 in migration of OS cells, 1x10⁶ MG-63 and U2OS cells were seeded in a 6-well plate, and 2 ml culture medium supplemented with 10% FBS were added. When cells were grown to 90% confluence, cells were synchronized in serum-free medium for 24 h, followed by a standard wound (<3 mm) created on the cell monolayer (time set as 0 h). A PBS solution was used to remove floating cells. Subsequently, cells were incubated in fresh complete medium (1% FBS) for 0, 24 and 48 h and the number of migrated cells was observed and counted under a light microscope. The results were quantified by Image-Pro plus software 6.0 (National Institutes of Health).

Transwell assay. The migration and invasion abilities of MG-63 and U2OS cells were measured using a transwell assay (cat. no. 3422; 6.5-mm insert; 8-µm polycarbonate membrane; Costar; Corning, Inc.). Briefly, 1x10⁶ transfected cells were suspended in serum-free medium and seeded into the upper chamber without (for the migration assay) Matrigel.
(BD Biosciences) or with a porous membrane coated with Matrigel (BD Biosciences) (for the transwell invasion assay). The Matrigel was incubated at 37˚C for 4 h before testing. Subsequently, serum-free medium was added into the upper chambers. Medium containing 10% FBS was added into the lower chambers. Following incubation for 48 h, the cells from the upper compartments were scraped off with cotton swabs, whereas the cells that migrated or invaded to the lower surface of the membrane were fixed with 4% paraformaldehyde at room temperature for 10 min and stained with 10% crystal violet at room temperature. The numbers of migrated or invasive cells were counted in five random fields under a light microscope at x200 magnification. Experiments were independently repeated three times.

Cell cycle and apoptosis analyses. The effect of lncGHET1 on cell cycle and apoptosis was evaluated by flow cytometry using the Annexin V Apoptosis Detection kit (BD Biosciences) and the Cell Cycle kit (BD Biosciences). Briefly, for cell cycle analysis, MG-63 and U2OS cells were harvested and washed with cold PBS twice. Following fixation with 70% ethanol at 4˚C overnight and labelling with propidium iodide (PI) solution (0.1 µg/µl) in the presence of Ribonuclease A (Takara Bio, Inc.) at 37˚C for 30 min in the dark, cells were assessed by flow cytometry (FACScan; BD Biosciences). For cell apoptosis analysis, cells were washed with cold PBS twice. Subsequently, cells were incubated with Annexin V-FITC and PI (BD Biosciences) for 15 min in the dark at room temperature. Finally, cell apoptosis was assessed using flow cytometry (FACScan; BD Biosciences) using FlowJo 10.06 software (FlowJo LLC). The data of these experiments were analyzed by FlowJo v10 software (Tree Star, Inc.).

Xenograft tumors in nude mice. Female BALB/c nude mice (SPF, n=6; age, 6 weeks; weight, 18-22 g) were provided by Nanjing Medical University (Nanjing, China). The mice were
housed in 12-h light/dark cycle in a temperature-controlled (22-26°C) and humidity-controlled (40-70%) room with freely accessible chow and tap water. To detect the effect of tumor growth in vivo, 5x10⁵ transfected MG-63 and U2OS cells were suspended in PBS and subcutaneously injected into the right flank of nude mice, and the mice of the control group were treated with PBS. Following a 36-day period, nude mice were euthanized by a high dose of anesthesia (150 mg/kg pentobarbital; intraperitoneal injection) according to protocols set by the Ethical Committee of the Ethical Committee of Nanjing Medical University (approval no. SYXK(SU)2018-0114). After nude mice were sacrificed, neoplasms were isolated for further analyses. The tumor volumes were recorded by slide calipers every week and calculated using the following formula: Volume=0.5 x length x width x width.

Hematoxylin and eosin (H&E) staining. The tumor tissues were fixed in 4% paraformaldehyde at room temperature for 10 min and then embedded in paraffin. The sections were immersed in 0.5% hematoxylin for 5 min, followed by staining with 0.5% eosin solution for 1 min at room temperature. At least three different sections of tumor tissues were examined for each group using a light microscope to assess the histopathological alterations.

Tunel assay. The in situ Cell Death Detection kit was purchased from Roche Diagnostics. All the steps were preformed according to the manufacturer's protocol. After washing with PBS for several times, sections (5-µm thick) were dewaxed in xylene and hydrated by graded ethanol solution. Samples were fixed in 10% formalin for 24 h at 4°C, embedded in paraffin and assessed using TUNEL staining (Roche Diagnostics GmbH) for 1 h at room temperature according to the manufacturer's protocol. After washing with PBS for several times, sections (5-µm thick) were incubated with 20 µg/ml proteinase K for 30 min at 37°C. Subsequently, the sections were incubated with a terminal deoxynucleotidyl transferase, followed by treatment with 3% hydrogen peroxide for 5 min. After that, the sections were incubated with peroxidase-conjugated antibody (cat. no. ab197034; dilution, 1:1,000; Abcam) for 10 min at room temperature. After washing with PBS for several times, the DAB solution with 3% hydrogen peroxide was added, and the methyl green was added. Finally, the sections were treated with Mayer's hematoxylin. The mounting medium was neutral resin. The images were observed in five fields under a fluorescence microscope.

Statistical analysis. All experiments were performed at least three times. The results are presented as the mean ± SD. Statistical analyses were performed using SPSS v19.0 software (IBM Corp.). Student’s t-test was performed to evaluate significant differences between two independent groups of samples. One-way ANOVA followed by Tukey’s post hoc test was applied to compare differences among multiple groups. P<0.05 was considered to indicate a statistically significant difference.

Results

lncGHET1 is upregulated in OS cell lines. The present study first examined the expression levels of IncGHET1 in the
U2OS, MG63, SaOs-2 and hFOB cell lines. Compared with the hFOB cell line, IncGHET1 expression was markedly higher in the OS cell lines (Fig. 1A). As shown in Fig. 1B, the cell proliferation abilities of U2OS, SaOs-2 and MG-63 cells were significantly higher than those in the hFOB group at 72 h. Furthermore, colony formation assays showed that there were more colonies in the former three cell lines compared with in the hFOB group (Fig. 1C). In addition, an increased migration and invasion rate was observed in the U2OS, SaOs-2 and MG-63 groups (Fig. 1D). These data indicated that IncGHET1 may function as a regulator of OS progression.

Effects of IncGHET1 on proliferation of OS cells. To assess the role of IncGHET1 in MG-63 and U2OS cell proliferation, siRNA was transfected to silence IncGHET1 expression. Following transfection for 48 h, IncGHET1 expression was detected using fluorescence microscopy and RT-qPCR. As shown in Fig. 2A and B, the results revealed that the siRNA transfection decreased IncGHET1 expression, whereas there was no significant difference observed between the si-NC and control groups. Subsequently, cell proliferation was assessed using CCK-8, colony formation and EdU assays. As demonstrated by the result of the CCK-8 assay, cell growth was suppressed in MG-63 and U2OS cells which were transfected with si-lncGHET1 compared with the cells transfected with si-NC and the control group (Fig. 2C). In addition, the colony formation ability of si-lncGHET1-transfected cells was decreased (Fig. 2D). An EdU assay was conducted to further confirm the effect of IncGHET1 on OS cells. The results revealed that downregulation of IncGHET1 markedly decreased the percentage of EdU-positive cells in MG-63 and U2OS cells (Fig. 3A).

To study the effect of IncGHET1 on the OS cell cycle, flow cytometry was performed. As shown in Fig. 3B, a significantly
Figure 4. Effect of lncGHET1 on cell growth in vivo. (A) Images of nude mice, (B) bodyweight, (C) tumor volume and (D) tumor weight are shown. The longest and shortest diameters of the tumor size in U2OS infected nude mice were 13.9, 20.1, 19.0 mm; 7.1, 13.4, 12.5 mm, successively. And these parameters of tumor size in the MG-63 infected nude mice were 15.1, 17.2, 20.0 mm; 10.9, 11.5, 11.3 mm, successively. (E) Hematoxylin and eosin staining was performed and the images were captured at a magnification of x200. \( *P<0.01 \) vs. control group. lncGHET1, long non-coding RNA gastric carcinoma proliferation enhancing transcript 1; NC, negative control; si, small interfering RNA.

Figure 5. Effect of lncGHET1 on cell apoptosis. (A) Apoptosis of tumors was detected using a TUNEL assay. Magnification, x200. (B) Apoptosis of MG-63 and U2OS cells was detected using flow cytometric analysis. (C) Key apoptotic players were examined by western blotting. lncGHET1, long non-coding RNA gastric carcinoma proliferation enhancing transcript 1; NC, negative control; si, small interfering RNA.
greater proportion of G1 phase cells and a markedly lesser proportion of S phase were observed in the si-lncGHET1 group compared with in the si-NC and control groups in both MG-63 and U2OS cells. Additionally, there was no significant difference among the three groups in G2 phase. Furthermore, western blot analysis was performed to determine the protein levels of relative factors, including cyclin D1, p21 and p27, involved in cell cycle regulation. As shown in Fig. 3C, compared with the si-NC and control groups, the expression levels of p21 and p27 protein were identified to be increased, whereas cyclin D1 expression was revealed to be decreased in the si-lncGHET1 group. Therefore, these data suggested that knockdown of lncGHET1 inhibited cell proliferation in MG-63 and U2OS cells.

Effect of lncGHET1 on development of xenograft tumors in vivo in OS. To further confirm the effect of lncGHET1 on the growth of tumors in vivo, MG-63 and U2OS cells transfected with si-lncGHET1 or si-NC were injected into nude mice. As shown in Fig. 4A-D, a significant decrease in tumor volume and weight was observed in the si-lncGHET1 group compared with in the control group and no significant change in body weight was observed. Additionally, H&E staining was performed to explore the histological alterations of tumors and the results demonstrated that the tissue structure was clearer and more complete in the si-lncGHET1 group than in the control group (Fig. 4E). These results suggested that silencing lncGHET1 expression inhibited the xenograft tumor growth of MG-63 and U2OS cells in vivo.

Effect of lncGHET1 on apoptosis of OS cells in vivo and in vitro. The present study investigated whether lncGHET1 knockdown affected cell apoptosis. The results of a Tunel assay revealed that the number of apoptotic cells was higher in nude mice in the si-lncGHET1 group compared with in mice in the control group (Fig. 5A). Flow cytometric analysis revealed that the percentage of apoptotic U2OS cells was notably increased in the si-lncGHET1 group, with 20.11% in the si-lncGHET1 group, 10.87% in the si-NC group and 9.11% in the control group of U2OS cells. Furthermore, a similar result was observed in MG-63 cells. A notable increase was identified in the si-lncGHET1 group (17.74%) compared with in the si-NC group (7.51%) and the control group (7.3%; Fig. 5B). Additionally, the present study examined several key apoptotic...
players in MG-63 and U2OS cells by western blot analysis (Fig. 5C). Compared with the control group, downregulation of lncGHET1 notably increased protein expression levels of Bax, caspase-3 and caspase-9, and decreased the expression levels of Bcl-2. These data indicated that reduced lncGHET1 expression promoted cell apoptosis in MG-63 and U2OS cells.

**Effect of lncGHET1 on cell migration and invasion in OS.** In order to verify the potential role of lncGHET1 in OS cell migration and invasion, wound healing, and transwell migration and invasion assays were conducted. The results of the wound healing assay revealed that cells migrated towards the wound at a slower rate in the si-lncGHET1 group compared with in the control group (Fig. 6A). In addition, transwell migration assay results indicated that knockdown of lncGHET1 markedly suppressed the migration ability of MG-63 and U2OS cells compared with that of cells in the control group, indicating a decreased migratory ability following lncGHET1
downregulation (Fig. 6B). Similarly, transwell invasion assays revealed that the number of invaded cells in the si-lncGHET1 group was lower than that in the control group (Fig. 6B).

Furthermore, western blotting was applied to evaluate the matrix metalloproteinase (MMP)-2 and MMP-9 protein levels. As shown in Fig. 6C, silencing lncGHET1 decreased the expression levels of MMP-2 and MMP-9 in MG-63 and U2OS cells compared with the control group. Therefore, the results suggested that the migration and invasion abilities of MG-63 and U2OS cells were suppressed following lncGHET1 knockdown.

Effect of lncGHET1 on cell EMT of OS. Increasing studies have demonstrated that EMT acts as an important step in metastatic dissemination of cancer cells (24-26). In the present study, the mRNA and protein expression levels of Snail, N-cadherin, E-cadherin and Vimentin were examined by RT-qPCR, western blotting and immunofluorescence analyses. The results revealed that E-cadherin mRNA expression was downregulated (Fig. 6B). Similarly, transwell invasion assays revealed that the number of invaded cells in the si-lncGHET1 group was lower than that in the control group (Fig. 6B).

Furthermore, western blotting was applied to evaluate the matrix metalloproteinase (MMP)-2 and MMP-9 protein levels. As shown in Fig. 6C, silencing lncGHET1 decreased the expression levels of MMP-2 and MMP-9 in MG-63 and U2OS cells compared with the control group. Therefore, the results suggested that the migration and invasion abilities of MG-63 and U2OS cells were suppressed following lncGHET1 knockdown.
significantly upregulated. The levels of protein expression were also increased, whereas the expression levels of Snail, N-cadherin and Vimentin were downregulated in MG-63 and U2OS cells which were transfected with si-lncGHET1 compared with in cells in the control group (Fig. 7A and B). As shown in Fig. 7C, compared with the si-NC and control group, knockdown of lncGHET1 in the MG-63 and U2OS cells was associated with lower expression of N-cadherin, while the expression levels of E-cadherin remained high.

**Knockdown of lncGHET1 inhibits cell proliferation and EMT via the Wnt/β-catenin signaling pathway.** Furthermore, the underlying mechanism of the role lncGHET1 serves in OS biology was explored. The present study investigated whether lncGHET1 could regulate the proliferation and EMT of OS cells via the Wnt/β-catenin signaling pathway. The present study silenced the expression of lncGHET1 in MG-63 and U2OS cells. Subsequently, the mRNA and protein expression levels of Wnt and β-catenin were detected using RT-qPCR, western blotting and immunofluorescence assays. The results demonstrated that expression levels of Wnt and β-catenin were decreased following lncGHET1 knockdown when compared with controls (Fig. 8). These data indicated that lncGHET1 exerted its effects in OS partly via the Wnt/β-catenin signaling pathway.

**Discussion**

OS is the most prevalent primary pediatric bone malignancy in the world (3,27-29). In recent years, advances of modern treatments have been used in the treatment of OS, such as multiagent chemotherapy, surgery and Chinese medical treatments (30,31). However, due to limitations of the effectiveness, the prognosis of OS is still unsatisfactory. The molecular mechanisms underlying the pathogenesis of OS have not been fully explored. Therefore, it is important to elucidate the predictive markers and underlying regulation process in OS.

In recent years, the roles of IncRNAs have attracted immense research interests worldwide. Aberrant expression of IncRNAs has been revealed to be associated with pathogenesis of cancer and other diseases (32-36). Previous studies have revealed that they serve important roles in physiological or pathology processes, particularly in cell growth, tumorigenesis, differentiation and development (15,37-42). However, to the best of our knowledge, the effect of lncGHET1 on the progression of OS remains undetermined.

To investigate the role of lncGHET1 in the regulation of OS, the present study first examined the expression levels of lncGHET1 in OS cell lines. The results revealed that all OS cell lines exhibited relatively high levels of lncGHET1 compared with hFOB cells. The results indicated that lncGHET1 may serve an important role in OS. In the present study, the functional role of lncGHET1 was analyzed in MG-63 and U2OS cells using loss-of-function approaches in vitro and in vivo. The results demonstrated that inhibition of lncGHET1 expression inhibited proliferation, migration and invasion, and suppressed cell cycle progression, whereas it promoted apoptosis of MG-63 and U2OS cells in vitro and in vivo, indicating lncGHET1 could be an oncogene in OS.

The EMT process contributes to the tumorigenesis (39). Increasing studies have demonstrated the vital role of EMT in cancer invasion and metastasis (25,43). It should be noted that EMT is characterized by loss of the epithelial characteristics, including loss of E-cadherin expression and upregulation of mesenchymal markers, such as vimentin and N-cadherin (44,45). Furthermore, a previous study has suggested that the EMT process is regulated by a set of transcription factors, including Snail, Slug and Twist (46). The present study suggested that suppressing lncGHET1 expression increased the expression levels of E-cadherin. However, Snail, N-cadherin and Vimentin expression was markedly decreased when lncGHET1 was silenced. Additionally, the present study explored the molecular mechanism by which lncGHET1 contributes to tumor progression, and indicated that lncGHET1 exerted an oncogenic effect, partly through regulating the Wnt/β-catenin signaling pathway (Fig. 9).

In conclusion, the present study demonstrated that inhibition of lncGHET1 attenuated cell proliferation, migration, invasion and EMT, and promoted apoptosis, partly through regulating the Wnt/β-catenin signaling pathway in OS cells. These findings indicated that lncGHET1 may serve as an oncogenic IncRNA in OS progression, and could be a promising molecular marker for diagnosis and treatment of OS.

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**Availability of data and materials**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Authors’ contributions**

WF designed the experiments. XC and WZ performed the experiments, and were the major contributors in writing the manuscript. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

All animal experiments were approved by the Ethical Committee of Nanjing Medical University.

**Patient consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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