Identification of \( \alpha_i3 \) as a promising target for osteosarcoma treatment

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Abstract

Sustained activation of multiple receptor tyrosine kinases (RTKs) simultaneously is vital for tumorigenesis and progression of osteosarcoma (OS). \( \alpha_i3 \) proteins recruitment to various RTKs mediates downstream oncogenic signaling activation. The expression, functions and underlying mechanisms of \( \alpha_i3 \) in human OS were examined. Expression of \( \alpha_i3 \) is robustly elevated in human OS tissues and is correlated with a poor overall survival. In patient-derived primary OS cells and immortalized lines (MG63 and U2OS), \( \alpha_i3 \) depletion, by shRNA and CRISPR/Cas9 strategies, robustly suppressed cell viability, proliferation and migration, while provoking G1-S arrest and apoptosis activation. Conversely, \( \alpha_i3 \) overexpressing ectopically can accelerate proliferation and migration of OS cells. In OS cells, \( \alpha_i3 \) immunoprecipitated with VEGFR2, FGFR, PGDFR and EGFR, mediating downstream cascade transduction. Akt-mTOR activation in primary OS cells was potently inhibited by \( \alpha_i3 \) shRNA, knockout or dominant negative mutation, but augmented after \( \alpha_i3 \) overexpression. In vivo studies showed that \( \alpha_i3 \) shRNA AAV (adeno-associated viruses) intratumoral injection largely inhibited the growth of subcutaneous xenografts of primary OS cells. Moreover, the growth of the \( \alpha_i3 \)-knockout primary OS xenografts was much slower than that of OS xenografts with empty vector. In \( \alpha_i3 \)-depleted OS xenografts tissues, \( \alpha_i3 \) downregulation and Akt-mTOR inactivation were detected. Taken together, overexpressed \( \alpha_i3 \) mediates RTK-Akt signaling to drive OS progression.

Key words: Osteosarcoma; \( \alpha_i3 \); multiple receptor tyrosine kinase; Akt-mTOR; Signaling

Introduction

For the advanced osteosarcoma (OS) patients with metastatic, recurrent or therapy-resistant OS, the prognosis is poor [1]. Further exploring the novel targeted therapeutics for OS is thus extremely important [2-7]. Due to various gene mutations, overexpression and/or over-activation of multiple receptor tyrosine kinases (RTKs) and its downstream oncogenic cascades are essential for initiation, progression and therapy-resistance of OS [4, 8, 9]. Several RTKs, including VEGFR, PDGFR, RET, EGFR and IGF2, as well as KIT and FGFR, are key drivers for the cancerous behaviors of OS [4, 8, 9]. Concurrent
activation of multiple RTKs shall provoke sustained activation of oncogenic cascades, causing persistent cancer growth [8, 9]. Therefore, targeting one or few RTKs, using genetic methods and/or pharmacological inhibitors, could only achieve minimal anti-OS efficiency [8, 9]. The novel strategies targeting multiple RTKs simultaneously should achieve better anti-OS outcome [8, 9].

The inhibitory guanine nucleotide regulatory proteins, Gαi proteins, consists of three subunits, including Gαi1, Gαi2 and Gαi3 [10]. It is known that GPCRs (G protein-coupled receptors) binding to Gαi proteins and the β and γ complexes will hinder adenylate cyclase, causing cyclic AMP (cAMP) depletion [10]. Such actions would be reversed by pertussis toxin [11]. Few studies have explored the expression, function and potential signaling mechanism of Gαi proteins in OS. Pine et al., have shown that Gαi proteins are important for the agonist-induced cAMP production in osteosarcoma cells that were derived from rat [12]. Wang et al., showed that pertussis toxin can inhibit bradykinin-induced Ca²⁺ mobilization in MG63 OS cells [13].

Our group has identified an essential role of Gαi proteins in transducing signals for multiple RTKs [14-20]. We found that EGFR-induced Akt-mTOR activation was abolished after Gαi3 double knockout (DKO) or silencing [20]. With VEGF stimulation, Gαi1/3 associated with VEGFR2, promoting VEGFR2 endocytosis and downstream signaling activation [16]. Similarly, Gαi1/3 mediated BDNF (brain-derived neurotrophic factor)-stimulated Akt-mTOR activation [17]. Therefore, by mediating signaling transduction of multiple RTKs, Gαi proteins could be important oncogenic genes and therapeutic targets for human cancer [14, 18, 21]. Indeed, we have previously shown that Gαi proteins are upregulated in glioma and gastric cancers, required for cancer growth [14, 18, 21]. The current study explored Gαi3 expression and potential functions in human OS.

**Methods**

**Ethics**

The protocols of the present study were reviewed and approved by the Ethics Committee of Soochow University and were in accordance with the principle of Helsinki declaration.

**Reagents**

LY294002 was from Sigma Aldrich (St Louis, M.O.). The antibodies were described in our previous studies [18]. All the primers, sequences, constructs and virus were synthesized by Shanghai Genechem Co. (Shanghai, China), or mentioned otherwise.

**Cells and tissues**

The immortalized OS lines, MG63 and U2OS, as well as the established hFOB1.19 osteoblastic cells were cultivated as described [22-24]. Patient-derived primary OS patients, namely pOS-1 and pOS-2, were described previously [22, 23]. Patient-derived human osteoblasts (pOsteoblasts) were differentiated and cultured using the previously-described protocols [25, 26]. The human tissues, including the OS tumor tissue specimens and the adjacent normal bone tissue specimens [22, 23], were obtained from the written-informed consent OS patients who were all administrated at the Affiliated Hospitals of Soochow University.

**Gene detection**

Protein detection by Western blotting, RNA assays by qRT-PCR and co-immunoprecipitation (co-IP) examining protein-protein interaction were extensively described early [17, 20]. When necessary, lysates from the same set of the experiment were detected in the parallel gels to test different proteins. The primers were described early [16].

**Gαi3 shRNA**

The lentivirus-encoded Gαi3 shRNAs and in vitro cell infection were described early [16]. Stable cells were selected by puromycin-containing complete medium (with FBS) for additional 96h. Gαi3 silencing (with over 95-98% knockdown efficiency) in the stable cells was always verified. The control cells were infected with non-sense scramble control shRNA lentivirus (‘sh-C’) [16]. For the in vivo studies, the Gαi3 shRNA sequence or shC sequence was sub-cloned into an established adenoviral vector, adeno-associated virus 9/AAV9 construct (Genechem). Through Lipofectamine 3000 the construct was thereafter transfected to HEK-293T cells, and the shRNA-expressing AAV virus was generated and was injected to xenograft tumors.

**CRISPR/Cas9-induced Gαi3 knockout (KO)**

OS cells were transfected with Cas9-expressing construct (Genechem) by Lipofectamine 3000 (Invitrogen, Shanghai, China) to establish stable cells. Next, the lenti-CRISPR/Cas-9 Gαi3 KO construct [14, 16], was transduced to Cas9-expressing OS cells, with stable cells established by using puromycin-containing medium for additional 96h. Gαi3 KO screening was carried out and thereafter the Gαi3 KO cells were eventually established. The control OS cells were with a lenti-CRISPR/Cas-9 empty vector with non-sense small guide RNA (“Cas9-C”).

**Gαi3 overexpression and dominant negative mutation**, stable cells establishment and verification
were described in our previous studies [14, 16].

**Constitutively-active mutant Akt1**

OS cells were infected with the constitutively-active Akt1 (caAkt1, S473D)-expressing adenovirus (provided by Dr. Li [27, 28]) for 48h. Puromycin was thereafter added for 96h to establish the stable OS cells, where expression of the caAkt1 was confirmed by Western blotting.

**Akt1/2 shRNA**

The commercial available Akt1/2 shRNA lentiviral particles (sc-43609-V, Santa Cruz Biotech) were added and transfected to cultured OS cells. After 48h, cells were cultured and selected in puromycin-containing medium for another 96h. Akt1/2 silencing was always examined.

**Cellular functional studies**

The cell viability detection by cell counting kit-8 (CCK-8), the EdU nuclear staining assay of cell proliferation, propidium iodide (PI)-FACS, “Transwell” assays were carried out using the previously described protocols [18, 21, 25, 29-33].

**Apoptosis detection**

Apoptosis-related assays, including the TUNEL nuclear staining, 7-AAD and Annexin V double staining, caspase-3/-9 activities measurement, and ELISA testing the cellular ssDNA (single strand DNA) contents were described early [21, 25, 31-33].

**Xenograft studies**

The primary pOS1 cells (five million cells in every mouse) were subcutaneously (s.c.) injected to the nude mice (18.5-19.5g, half female and half male, please refer to our previous studies [22, 23]). Tumor-bearing mice were then subject to the designated treatments. Tumor volumes \((\text{length} \times \text{width}^2)/2\) and animal body weights were weekly recorded. Soochow University’s Ethics Committee and Institutional Animal Care and Use Committee (IACUC) reviewed and approved the protocols of animal studies.

**Statistical analyses**

Statistical analyses were described previously [22, 23]. The numerical data in the bar graphs indicated the mean and standard deviation (S.D.). \(P\)-values < 0.05 were statistically significant.

**Results**

**In human OS Gαi3 is upregulated**

The Cancer Genome Atlas (TCGA) database (available on the public domain https://portal.gdc.cancer.gov) was first consulted to analyze Gαi3 transcripts in human sarcoma tissues. Total 264 samples (HTSeq-FPKM) were collected, including the two normal specimens and the 262 sarcoma specimens. As shown, Gαi3 transcripts in the retrieved sarcoma tissue specimens were higher than those in the retrieved normal tissue specimens (Figure 1A). The Kaplan-Meier survival, Figure 1B, verified that high Gαi3 expression in sarcoma patients was correlated with a poor prognosis (HR = 1.73, \(P = 0.008\)). Subgroup analysis by different clinical features demonstrated that Gαi3-high expression was associated with a poor prognosis in male sarcoma patients (\(P = 0.003\), Figure 1C), age≤60 (\(P = 0.048\), Figure 1D), Tumor depth: deep (\(P = 0.009\), Figure 1E).

Next, we tested Gαi3 expression in local OS tissues. As previously described [22, 23], OS tissue specimens and matched adjacent bone normal tissue specimens were retrieved from 10 different OS patients. Figure 1F showed that Gαi3 mRNA in the OS tissue specimens (“T”) was more than six-fold higher than that in the adjacent normal bone tissues (“N”). Gαi3 protein expression was tested as well. In the OS tissues of four representative OS patients (Patient #1/#2/#3/#4), protein expression of Gαi3 was significantly elevated (Figure 1G). When combining all blotting results of the ten sets tissues, Gαi3 protein was found to be significantly upregulated in the OS tissues (\(P < 0.001\) versus “N” tissues, Figure 1H). The immunohistochemistry (IHC) staining assay results, in Figure 1I, verified Gαi3 protein elevation in OS tumor tissues (Figure 1I). While low expression of Gαi3 was detected in the adjacent normal tissue specimens (Figure 1I).

The expression of Gαi3 in different human OS cells was examined. Patient-derived primary OS cells (pOS-1/2, from two different OS patients), as well as the immortalized cells (MG63 and U2OS lines), were cultured. The expression of Gαi3 mRNA in different primary and immortalized OS cells was dramatically higher than that in the hFOB1.19 human osteoblastic cells and patient-derived human osteoblasts (“pOsteoblasts”) (Figure 1J). In addition, Gαi3 protein upregulation was also shown in different primary and established OS cells (Figure 1K). Taken together, in OS tissues and cells Gαi3 is upregulated.

**Gαi3 silencing exerts anti-tumorigenic activity in cultured OS cells**

pOS-1 primary cells were separately transfected with two different Gαi3 lentiviral shRNAs, sh-Gαi3- seq1/sh-Gαi3-seq2 [22, 23]. After puromycin-induced selection, Gαi3-silenced stable OS cells were established. Gαi3 mRNA expression decreased over 90-95% in sh-Gαi3-expressing stable pOS-1 cells
Expression of Gαi1 and Gαi2 mRNA was unaffected (Figure 2A). Gαi3 protein silencing was detected in the stable pOS-1 cells with Gαi3 shRNAs. Gαi1 and Gαi2 protein expression was again unchanged (Figure 2B).

CCK-8 assays showed that Gαi3 shRNA potently decreased pOS-1 cell viability (CCK-8 OD, Figure 2C). Moreover, Gαi3 shRNA potently inhibited pOS-1 cell proliferation, as the ratio of EdU positively stained nuclei was robustly decreased in sh-Gαi3-expressing pOS-1 cells (Figure 2D). The PI-FACS assays were employed to test cell cycle progression. The ratio of G1-phase cells was significantly increased in pOS-1 cells expressing Gαi3 shRNA (Figure 2E), where S-phase pOS-1 cell percentage was decreased (Figure 2F). These results implied that the designated Gαi3 shRNAs induced G1-S arrest in primary OS cells. The “Transwell” results demonstrated that Gαi3 shRNA dramatically inhibited pOS-1 cell in vitro migration (Figure 2G). The scramble control shRNA, or shC, did not significantly alter Gαi1/2/3 expression (Figure 2A-B) and affect OS cellular behaviors (Figure 2C-G).

Figure 1. In human OS Gαi3 is upregulated. TCGA cohorts show Gαi3 mRNA transcripts in 262 sarcoma cases (“Primary Tumor”) and two normal tissue cases (A). The Kaplan Meier Survival curve of Gαi3-low (n = 130, blue) and Gαi3-high (n = 132, red) sarcoma patients was presented (B). Subgroup analyses, based on the different clinical features of the OS patients, were performed as well (C-E). The expression of Gαi3 (both mRNA and protein) in twelve (n = 12) pairs of OS tumor tissue specimens (“T”) and adjacent normal bone tissue specimens (“N”) was tested (F-H). The representative human tissue Gαi3 IHC were presented as well (I). The expression of Gαi3 mRNA and protein in the mentioned cells was measured (J-K). *P < 0.05 versus “N” tissues (F and H) or hFOB1.19 cells (J and K). Scale bar = 100 µm (I).
Next, OS cells that were derived from another OS patient, pOS2 [22, 23], and the immortalized lines (MG63 and U2OS), were stably transduced with the lentiviral Gαi3 shRNA (sh-Gαi3-seq1, “sh-Gαi3”). As shown, the designated shRNA resulted in robust Gαi3 mRNA downregulation in both primary and immortalized OS cells (Figure 2H). Gαi3 shRNA largely suppressed CCK-8 viability (Figure 2I), proliferation (the ratio of EdU positively stained nuclei reduction, Figure 2J) and migration (Figure 2K) in pOS2 primary cells and established lines.

The potential effect of Gαi3 shRNA on the Gαi3-low human osteoblasts (see Figure 1J-K) was tested as well. hFOB1.19 osteoblastic cell line or patient-derived primary osteoblasts (pOsteoblasts) were stably transduced with the lentiviral sh-Gαi3-seq1, causing dramatic Gαi3 mRNA downregulation (“sh-Gαi3”, Figure 2L). Interestingly, in hFOB1.19 cells and pOsteoblasts, Gαi3 shRNA failed to significantly inhibit CCK-8 viability (Figure 2M) and cell proliferation (by measuring EdU positively stained nuclei ratio, Figure 2N), supporting a cancer cell-specific effect by Gαi3 shRNA.
Ga\(\alpha\)i3 silencing provokes apoptosis activation in OS cells

Since Ga\(\alpha\)i3 silencing induced viability reduction, growth arrest, proliferation inhibition, we therefore tested its potential function on apoptosis in OS cells. Results showed that the relative activities of caspase-3 and caspase-9 (Figure 3A and B) were augmented in sh-Ga\(\alpha\)i3-expressing stable pOS-1 cells. Figure 3C showed that Ga\(\alpha\)i3 shRNA induced cleavages of caspase-3, PARP and caspase-9 in pOS-1 primary cells. Confirming increased DNA breaks in pOS-1 cells with Ga\(\alpha\)i3 shRNA, we showed that ssDNA contents were dramatically increased (tested by the ELISA assays, Figure 3D). Moreover, FACS assay results in Figure 3E showed that Ga\(\alpha\)i3 silencing increased the pOS-1 cell number with Annexin V-7-AAD double positive staining, confirming apoptosis activation. As expected, shC did not provoke caspase-apoptosis activation in the primary pOS-1 cells (Figure 3A-E).

In pOS-2 cells and immortalized lines (MG63 and U2OS), Ga\(\alpha\)i3 shRNA (sh-Ga\(\alpha\)i3-seq1, “sh-Ga\(\alpha\)i3”, see Figure 2) similarly augmented the relative caspase-3 activity (Figure 3F). Apoptosis induction was observed as well in the sh-Ga\(\alpha\)i3-seq1-expressing pOS-2 cells and immortalized cell lines, evidenced by the significantly increased Annexin V-7-AAD double staining (Figure 3G). Conversely, in the osteoblastic cell line hFOB1.19 and pOsteoblasts, shRNA-induced silencing Ga\(\alpha\)i3 (“sh-Ga\(\alpha\)i3”, see Figure 2) failed to significantly induce apoptosis (by quantifying TUNEL-positively stained nuclei, Figure 3H).

Ga\(\alpha\)i3 knockout potently inhibits OS cell progression in vitro

Next, a previously-described CRISPR/Cas9-Ga\(\alpha\)i3-KO-puro construct [15, 16] was transduced to the Cas9-expressing pOS-1 primary cells. Thereafter, single stable pOS-1 cells with the Ga\(\alpha\)i3 KO construct, or the ko-Ga\(\alpha\)i3 cells, were established after Ga\(\alpha\)i3 KO screening, and Ga\(\alpha\)i3 mRNA and protein (Figure 4A) depletion was detected. Ga\(\alpha\)i3 KO largely decreased CCK-8 viability (Figure 4B) and inhibited cell proliferation (the ratio of EdU positively stained nuclei decreasing, Figure 4B) and migration (Figure 4C) in pOS-1 cells. On the contrast, caspase-3 and caspase-9 activities (Figure 4D) were augmented in ko-Ga\(\alpha\)i3 pOS-1 cells, where caspase-3, PARP and caspase-9 cleavages were induced (Figure 4D). In addition, apoptosis induction was detected in the ko-Ga\(\alpha\)i3 pOS-1 cells, as the ratio of TUNEL-positively stained nuclei decreasing, Figure 4B) and inhibited cell proliferation (the ratio of EdU positively stained nuclei decreasing, Figure 4B) and migration (Figure 4C) in pOS-1 cells.
stained nuclei was increased significantly (see the quantified results in Figure 4E).

The Gai3-KO construct was employed to knockout Gai3 in primary pOS-2 cells and immortalized lines (MG63 and U2OS), and stable cells established (“ko-Gai3”) after screening (Figure 4F). As shown, Gai3 KO inhibited cell proliferation (the ratio of EdU positively stained nuclei reduction, Figure 4G) and in vitro migration (Figure 4H) in pOS-2 and established OS cells. Increased TUNEL-positive nuclei ratio confirmed apoptosis activation in ko-Gai3 pOS-2 cells and immortalized lines (Figure 4I). Together, Gai3 KO, by the CRISPR/Cas9 strategy, resulted in profound anti-OS cell activity.

Further promoting OS cell growth by Gai3 ectopic overexpression

Next, a Gai3-expressing lentiviral construct (see our previous studies [14, 16]) was transduced to pOS-1 cells. Following selection Gai3-overexpressed stable pOS-1 cells were thereafter established: namely “OE-Gai3-sL1” and “OE-Gai3-sL2” (two selections). Gai3 mRNA and protein expression levels (Figure 5A and B) were robustly elevated in the OE-Gai3 pOS-1 cells. Gai3 overexpression accelerated pOS-1 cell proliferation (the ratio of EdU positively stained nuclei increasing, Figure 5C) and in vitro migration (“Transwell” assays, results quantified in Figure 5D).

In pOS-2 cells and the immortalized lines (MG63 and U2OS), ectopic overexpression of Gai3 using the same construct (“OE-Gai3”, Figure 5E) enhanced cell proliferation (the ratio of EdU positively stained nuclei increasing, Figure 5F) and in vitro migration (see the quantified results in Figure 5G). Therefore, these results again supported the key cancer-promoting function of Gai3 in OS cells.

Gai3 immunoprecipitates with RTKs and is key to Akt-mTOR activation in OS cells

Our previous studies have shown that Gai proteins associated with several oncogenic RTKs (EGFR, VEGFR2, TrkB, FGFR and KGFR), mediating downstream signaling activation [14, 16-21]. Co-IP (co-immunoprecipitation) assays, Figure 6A, demonstrated that Gai3 immunoprecipitated with VEGFR2, FGFR, PGDFR and EGFR in primary OS cells (“pOS-1/2”) and immortalized U2OS cells. Moreover, the association between Gai3 and multiple RTKs (VEGFR2, FGFR, PGDFR and EGFR) was detected in the human OS tissues from three representative patients (Figure 6B). When testing downstream Akt-mTOR signaling, we showed that levels of phosphorylated-Akt (Ser-473) and phosphorylated-S6K (Thr-389) were dramatically decreased in pOS-1 cells bearing Gai3 shRNAs (Figure 6C). Moreover, CRISPR/Cas9-induced Gai3

Figure 4. Gai3 knockout potently inhibits OS cell progression in vitro. Patient-derived OS cells (“pOS-1/2”) or immortalized OS lines (MG63 and U2OS), bearing the CRISPR/Cas9-Gai3-KO-puro construct (“ko-Gai3”) or the corresponding vector (“Cas9-C”), were established, expression of listed mRNAs and proteins were examined (A and F); After culturing for the designated hours, CCK-8 viability (B), proliferation (testing the ratio of EdU positively stained nuclei, B and G), cell migration (“Transwell” assays, C and H), caspase-PARP activation was tested (D), with cell apoptosis measured through quantifying the TUNEL-positively stained nuclei ratio (E and I). All blotting data in this Figure were repeated five times. *P < 0.05 versus “Cas9-C” group. Each single experiment was repeated for five times. Scale bar = 100 µm (B and C).
KO (see Figure 4) largely inhibited Akt-S6K phosphorylations in pOS-1 cells (Figure 6D). Notably, RTKs (FGFR, PGDFR and EGFR) expression and phosphorylation were unaffected by Gαi3 shRNA (sh-Gαi3-seq1) or Gαi3 KO (Figure 6D). Conversely, ectopic overexpression of Gαi3 (see Figure 5) significantly increased Akt-S6K activation in pOS-1 cells (Figure 6E).

Figure 5. Further promoting OS cell growth by Gαi3 ectopic overexpression. Patient-derived primary OS cells (“pOS-1/-2”) or immortalized OS lines (MG63 and U2OS), bearing the lentiviral construct encoding wild-type Gαi3 (“OE-Gαi3”) or the corresponding vector (“Vec”), were established and cultivated, expression of listed mRNAs and proteins were measured (A, B and E); After culturing for the designated hours, cell proliferation (testing the ratio of EdU positively stained nuclei, C and F) and cell migration (“Transwell” assays, D and G) were measured. All blotting data in this Figure were repeated five times. *P < 0.05 versus “Vec” group. Each single experiment was repeated for five times. Scale bar = 100 µm (C and F).

Figure 6. Gαi3 immunoprecipitates with RTKs and is key to Akt-mTOR activation in OS cells. The association between Gαi3 and the designated RTKs (VEGFR2, FGFR, PGDFR and EGFR) in patient-derived primary human OS cells (“pOS-1/-2”) and U2OS line (A, cultured in FBS-containing medium for 5 min) as well as in OS tissues of the representative patients (B) was examined by the co-immunoprecipitation (Co-IP) assays. The pOS-1 primary cells stably expressing the Gαi3 shRNA (sh-Gαi3-seq1/sh-Gαi3-seq2), the CRISPR/Cas9-Gαi3-KO-puro construct (“ko-Gαi3”), the Gαi3-expressing lentiviral construct (“OE-Gαi3”), or their corresponding controls (“shC”, “Cas9-C” or “Vec”) were established, and expression of listed proteins tested (C-E). The pOS-1 primary cells, stably expressing the lentiviral dominant negative Gαi3 construct (dnGαi3) or the empty vector (“Vec”), were established, the association between Gαi3 and RTKs (VEGFR2, FGFR, PGDFR and EGFR) as well as their expression were examined (F and G). After culturing for the designated hours, cell proliferation and migration were separately examined by EdU staining (H) and “Transwell” (I) assays. All blotting data in this Figure were repeated five times. “pare” indicated the parental control OS cells. *P < 0.05 versus “shC”/“Cas9-C”/“Vec” group. Each single experiment was repeated for five times.
Figure 7. Akt-mTOR inhibition contributes to Gαi3 depletion-induced anti-OS cell activity. The pOS-1 cells bearing the CRISPR/Cas9-Gαi3-KO-puro construct (“ko-Gαi3”) were further infected with the constitutively-active Akt1 adenovirus (“Ad-caAkt1”) or the adenovirus with the empty vector (“Ad-Vec”), control cells were with the CRISPR/Cas9 empty vector (“Cas9-C”), listed proteins were shown (A). Cells were cultured for designated hours, cell proliferation (B, EdU assays) and migration (C, “Transwell” assays) were tested. pOS-1 cells stably bearing the lentiviral Akt1/2 shRNA (“shAkt1/2”) were further transduced with a wild-type Gα3 (“OE-Gα3”) lentiviral construct, the lentiviral sh-Gαi3-seq1 (“sh-Gαi3”) or their control construct (“Vec+shC”), stable cells were established. Gαi3 mRNA and listed proteins were shown (D). After culturing for the designated hours, cell proliferation (E, by measuring EdU positively stained nuclei ratio) and migration (F) were measured. pOS-1 cells, bearing the lentiviral construct encoding wild-type Gαi3 (“OE-Gαi3-sL1”) were treated with LY294002 (150 nM) or the vehicle control (0.1% DMSO), and cultured for designated time periods, listed proteins were shown (H), with cell proliferation (I) and migration (J) examined as well. All blotting data in this Figure were repeated five times. “pare” indicated the parental control OS cells. *P < 0.05 (A-C). *P < 0.05 versus “pare” cells (E-G). *P < 0.05 versus “DMSO” (I and J). “N. S.” indicated no statistical difference (P > 0.05, E and F). Each single experiment was repeated for five times. Scale bar = 100 µm (B, C, E, I and J).

To block Gαi3-RTKs association, the lentiviral dominant negative (dn) Gαi3 mutant construct was stably transduced into pOS-1 cells. The dnGαi3 mutant will replace the conserved Gly (G) residue with the Thr (T) residue in the G3 box, thereby preventing Gαi1/3 interaction with the associated proteins [19, 20]. Results show that dnGαi3 disrupted the association between Gαi3 and multiple RTKs (VEGFR2, FGFR, PDGFR and EGFR) in pOS-1 cells (Figure 6F). Expression of RTKs was however unchanged (Figure 6F, “Input”). Importantly, dnGαi3 largely inhibited Akt-S6K phosphorylations in pOS-1 primary cells (Figure 6G). The dnGαi3 largely suppressed pOS-1 cell in vitro proliferation and migration, examined through the nuclear EdU staining (see the quantified results in Figure 6H) and “Transwell” (Figure 6I) assays, respectively.

Akt-mTOR inhibition contributes to Gαi3 depletion-induced anti-OS cell activity

To support that Akt-mTOR inhibition was the main mechanism of Gαi3 depletion-caused anti-OS cell activity, we expressed the constitutively active Akt1 (caAkt1) [34] adenovirus (“Ad-caAkt1”) that could rescue Akt and S6K phosphorylation in koGαi3 pOS-1 cells (Figure 7A). Significantly, Ad-caAkt1 restored proliferation (by quantifying EdU-positively stained nuclei, Figure 7B) and in vitro migration

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“Transwell” assays, Figure 7C) of koGαi3 pOS-1 cells. Furthermore, shRNA-induced silencing of Akt1/2 (Figure 7D) blocked Akt-S6K phosphorylations (Figure 7D) and mimicked Gαi3 depletion-induced actions, suppressing pOS-1 cell proliferation (Figure 7E) and in vitro migration (Figure 7F). Significantly, re-introducing the Gαi3 shRNA lentivirus or the Gαi3-expressing construct (Figure 7G) was unable to further influence cell proliferation (Figure 7F) and migration (see quantified results in Figure 7F) in Akt1/2-silenced pOS-1 cells. Moreover, in the Gαi3-overexpressed pOS-1 cells (OE-Gαi3-sL1), treatment with LY294002, a PI3K-Akt inhibitor [35], blocked Akt-S6K phosphorylation (Figure 7H) and inhibited cell proliferation (Figure 7I) and in vitro cell migration (Figure 7J). Thus, Akt-mTOR inhibition should be responsible for Gαi3 depletion-induced anti-OS cell activity.

Gαi3 depletion inhibits OS cell growth in vivo

At last, pOS-1 cells were subcutaneously (s.c.) injected to the nude mice. Within 20 days of cell inoculation, the subcutaneous pOS-1 xenograft tumors were established and each tumor volume was close to 100 mm³ (“Day-0”). The xenograft-bearing mice were thereafter assigned into three different groups randomly, with six mice in every group (n = 6). Afterwards, the mice were intratumorally injected daily with the AAV-packed Gαi3 shRNAs (AAV-sh-Gαi3-seq1 or AAV-sh-Gαi3-seq2, two different sequences in the AAV9 construct) or AAV-packed control shRNA (AAV-shC), for 12 consecutive days. Figure 8A, recording tumor growth, demonstrated that the growth of pOS-1 xenograft tumors was robustly mitigated after AAV-sh-Gαi3 injection. The volumes of AAV-sh-Gαi3-injected pOS-1 xenografts were dramatically lower than those of AAV-shC-injected pOS-1 xenografts (Figure 8A). The estimated daily tumor growth (see previous studies [22, 23]) results demonstrated that subcutaneous pOS-1 xenograft growth was dramatically suppressed with AAV-sh-Gαi3 injection in the nude mice (Figure 8B). At Day-42, all the animals were anaesthetized and decapitated, and pOS-1 xenografts carefully isolated and weighted. The pOS-1 tumors with AAV-sh-Gαi3 injection were significantly lighter than pOS-1 tumors with the control shRNA virus injection (Figure 8C). Among the three mice groups there was no significant difference in the mice body weights (Figure 8D).

![Figure 8. Gαi3 depletion inhibits OS cell growth in vivo.](https://www.ijbs.com)
At experimental Day-7 and Day-14, 3h after virus injection, one tumor of each group was carefully isolated, and total six tumors were obtained. Ga\textsubscript{i3} mRNA was dramatically decreased in AAV-sh-Gai3-injected pOS-1 xenograft tissues (Figure 8E), where Ga\textsubscript{i3} protein downregulation as well as p-Akt and p-S6K inhibition were detected (Figure 8F). Supporting apoptosis activation, we showed that cleaved-caspase-3/cleaved-PARP levels were augmented in Ga\textsubscript{i3}-silenced pOS-1 xenograft tissues (Figure 8G). Thus, intratumoral injection of AAV-packed Ga\textsubscript{i3} shRNA suppressed Akt-mTOR activation and provoked apoptosis in pOS-1 xenografts.

In addition the ko-Ga\textsubscript{i3} pOS-1 cells and the Ca9-C control cells (see Figure 4) were injected to the nude mice, forming subcutaneous xenografts. After 20 days, tumor recordings were started (“Day-0”). As shown ko-Ga\textsubscript{i3} pOS-1 xenograft growth was slower than the Ca9-C xenografts (Figure 8H), while animal body weights were indifferent (Figure 8I). At experimental Day-7, we carefully isolated one tumor xenograft per group. Ga\textsubscript{i3} mRNA and protein (Figure 8J-K) expression was completely depleted in ko-Ga\textsubscript{i3} pOS-1 xenograft tissues, where p-Akt was decreased (Figure 8K). The cleaved-caspase-3 and the cleaved-PARP levels were increased in Ga\textsubscript{i3}-KO xenograft tissues (Figure 8K), supporting apoptosis induction in vivo.

**Discussion**

The GPCR superfamily is composed of the immense structural and functional different proteins, participating in various biological processes and signals in the bone [36]. Due to gene mutations, depletion or overexpression, GPCR components are dysregulated in human OS [36]. Their roles in OS progression have been established [36]. For example, Iyer et al., found that A3 adenosine receptor (A3AR) depletion activated protein kinase A (PKA)–Akt–nuclear factor (NF)-\kappaB signaling to promote OS cell growth [37]. High GPR56 (G protein-coupled receptor 56) expression is an unfavorable prognostic factor, promoting invasion and proliferation of OS cells [38]. Liu et al. demonstrated that GPR110 (G protein-coupled receptor 110) silencing inhibited OS cell growth [39].

Importantly, studies have reported that Ga\textsubscript{i1/3} coupled GPCRs, including Apelin receptors [40], CXCR4 [41-43], melatonin receptors [44], are important contributor for OS progression. Due to various gene mutations (or overexpression), concurrent and sustained activation of multiple different RTKs in OS will provoke sustained activation of oncogenic signaling, leading to persistent OS growth and progression [4, 8, 9]. Interestingly, we have previously shown that Ga\textsubscript{i} proteins are essential for signalings by several important oncogenic RTKs (including EGFR [20], VEGF2 [16], KGF [19], FGFR [18] and TrkB [17]) as well as the non-RTK receptor (IL-4R) [15].

After showing the essential role of Ga\textsubscript{1/3} in activation of oncogenic signalings by RTKs, we previously explored Ga\textsubscript{1/3} in different human cancers. Ga\textsubscript{1} and Ga\textsubscript{3} expression is elevated in glioma, correlating with tumor stage [14, 18]. Ga\textsubscript{1} can form a complex with multiple RTKs (including FGFR, PDGFR and EGFR), transducing downstream Akt-mTOR activation in glioma tissues and cells [18]. Conversely, Ga\textsubscript{i1} silencing or mutation inhibited glioma cell growth [18]. In the mouse brain, the orthotopic growth of patient-derived glioma xenografts was arrested after Ga\textsubscript{1/3} depletion, whereas forced overexpression of Ga\textsubscript{1/3} enhanced growth [14]. We also showed that Ga\textsubscript{1} upregulation in human gastric cancer was correlated with poor overall survival [21]. Ga\textsubscript{1} silencing or knockout inhibited Akt-mTOR activation and gastric cancer cell growth [21]. These previous studies supported that Ga\textsubscript{1/3} could be important oncogenic genes and promising therapeutic targets of human cancer.

Ga\textsubscript{i3} should be a vital gene for OS progression. TCGA database shows that transcripts of Ga\textsubscript{i3} are significantly upregulated in sarcoma tissues, and high-Ga\textsubscript{i3} expression in sarcoma correlating with the poor overall survival. Ga\textsubscript{i3} elevation was observed in local OS tissues as well as in different immortalized and primary OS cells, while low expression was observed in cancer-surrounding normal bone tissues and in immortalized and primary osteoblasts. Functional studies showed that in different OS cells, Ga\textsubscript{i3} depletion, by shRNA or CRISPR/Cas9 strategies, robustly suppressed cell survival, proliferation and cell migration, and provoking G1-S arrest and apoptosis. Contrarily, ectopic Ga\textsubscript{i3} overexpression can further accelerate OS cell growth. In vivo, Ga\textsubscript{i3} shRNA AAV intratumoral injection potently suppressed the growth of the patient-derived OS xenografts in nude mice. Moreover, the growth of primary OS xenografts of the Ga\textsubscript{i3} KO cells was largely suppressed.

We have previously discovered that Ga\textsubscript{i1/3} association with multiple RTKs was required for downstream signaling activation. For instance, Ga\textsubscript{i1/3} are key proteins in mediating VEGF-induced VEGFR2 signaling [16]. Following VEGF stimulation, Ga\textsubscript{i1/3} were in the VEGFR2 endocytosis complex, required for VEGFR2 endocytosis and subsequent activation of downstream signalings [16]. Similarly, Ga\textsubscript{i1/3} proteins are indispensable signaling molecule.
for EGF- and KGF-induced Akt-mTORC1 signaling activation [19, 20]. In addition, brain-derived neurotrophic factor (BDNF)-induced signaling and anti-depressive actions required Gαi1/3 [17]. Gαi 1/3 silencing inhibited BDNF-induced TrkB endocytosis and activation of the downstream signaling [17].

The present study implied that Gαi3-driven OS cell growth was primarily through mediating Akt-mTOR cascade activation. In OS cell and tissues Gαi3 associated with RTKs (VEGFR2, FGFR, PGDFR and EGFR), essential for downstream Akt-mTOR activation. In OS cells Akt-S6K activation was largely inhibited Akt-S6K activation and OS cell proliferation and migration. Restoring Akt-S6K expression failed to affect proliferation and migration in Akt1/2-silenced cells. Therefore, Gαi3-silencing inhibited OS cell proliferation and depletion-induced actions, Akt1/2-depleted OS xenograft tissues. Importantly, Gαi3-associated with RTKs (VEGFR2, FGFR, PGDFR and EGFR), essential for downstream Akt-mTOR cascade activation. In OS cell and tissues Akt-S6K phosphorylations were decreased in Gαi3-depleted OS xenografts tissues. Importantly, disrupting Gαi3-RTKs association, through dnGαi3, largely inhibited Akt-S6K activation and OS cell proliferation and migration. Restoring Akt-S6K activation, by caAkt1, rescued proliferation and migration of Gαi3-KO OS cells. Conversely, mimicking Gαi3 deletion-induced actions, Akt1/2 silencing inhibited OS cell proliferation and migration. Significantly, exogenously altering Gαi3 expression failed to affect proliferation and migration in Akt1/2-silenced cells. Therefore, Gαi3-driven OS cell growth was possibly due to mediating RTKs-Akt signaling.

Conclusion

Over three-fifths of bone sarcoma are OS [45, 46]. The standard chemotherapy of OS in clinic is the combination of methotrexate, doxorubicin, and cisplatin [45, 47, 48], showing limited success in metastatic and other OS patients with advanced diseases [45, 47, 48]. Further exploring key pathologic mechanisms and the driving signaling molecule for advanced OS is therefore important [45, 47, 48]. The results of this study showed that overexpressed Gαi3 mediated RTKs signaling to drive OS progression, serving as a novel and promising treatment molecular target for patients with OS.

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Ethics Statement

This study was approved by Ethics Committee of Soochow University.

Data Availability Statement

All data are available upon request.

Author Contributions

All the listed authors conceived, designed, and supervised the study. All listed authors collected samples, performed the experiments and analyzed the data. All authors reviewed and approved the final manuscript.

Competing Interests

The authors have declared that no competing interest exists.

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