Downregulation of TLX induces TET3 expression and inhibits glioblastoma stem cell self-renewal and tumorigensis

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Glioblastomas have been proposed to be maintained by highly tumorigenic glioblastoma stem cells (GSCs) that are resistant to current therapy. Therefore, targeting GSCs is critical for developing effective therapies for glioblastoma. In this study, we identify the regulatory cascade of the nuclear receptor TLX and the DNA hydroxylase Ten eleven translocation 3 (TET3) as a target for human GSCs. We show that knockdown of TLX expression inhibits human GSC tumorigenicity in mice. Treatment of human GSC-grafted mice with viral vector-delivered TLX shRNA or nanovector-delivered TLX siRNA inhibits tumour development and prolongs survival. Moreover, we identify TET3 as a potent tumour suppressor downstream of TLX to regulate the growth and self-renewal in GSCs. This study identifies the TLX-TET3 axis as a potential therapeutic target for glioblastoma.
Glioblastoma (GBM) is the most common and aggressive primary brain tumour with median survival time of 14 months after diagnosis. No effective treatment has been developed for GBM patients yet. It has been proposed that GBMs are maintained by a small population of cancer stem cells that retain stem cell properties, are highly tumorigenic and resistant to radiotherapy and chemotherapy. The cancer stem cell hypothesis proposes cancer stem cells reside at the top of a cellular hierarchy and have the ability to give rise to the heterogeneous populations of the tumour bulk. The presence of cancer stem cells together with the heterogeneity of the tumour mass renders GBM treatment resistant and recurring. Therefore, new therapies are needed to target these cancer stem cells.

TLX (NR2E1) is a nuclear receptor expressed in vertebrate forebrains and essential for neural stem cell self-renewal. Recently, TLX has been shown to be expressed in human GBM tissues and cell lines and play a role in GBM development in mouse tumour models. However, the function of TLX in human glioblastoma stem cell (GSC)-initiated tumorigenesis and the effect of modulating TLX expression in human GSCs on the development of GBM remain to be determined.

5-Hydroxymethylcytosine (5hmC) is a form of DNA modification derived from hydroxylated 5-methylcytosine (5mC). The level of 5hmC is considerably reduced in many types of human cancers, including gliomas. 5hmC is tightly controlled by the TET family of dioxygenases, which catalyse the conversion of 5mC to 5hmC. TET proteins have been shown to inhibit haematopoietic transformation, breast and prostate cancer invasion and metastasis. However, the role of TET proteins, especially TET3, in GBM tumorigenesis remains largely unknown.

RNA interference holds great promise for tumour therapy. However, efficient delivery of small RNAs in vivo represents a major challenge preventing RNA interference from achieving the potency required for successful clinical applications. After ups and downs, RNA interference is now regaining its momentum. Various delivery technologies have been developed for RNA interference. Viral vectors have high delivery efficiency and allow sustained gene silencing with a single injection, offering practical advantage for diseases associated with hard-to-reach organs, such as the brain. Non-viral vectors, such as cationic lipids and polymers, are developed to increase safety and efficiency of delivery.

Dendrimers are one of the most promising non-viral vectors for delivering small RNAs by virtue of their well-defined structure and unique multivalent cooperativity alongside the high payload confined within a nanosized volume. In particular, ploy(amideamine) dendrimers bear amine groups at the terminals, which can effectively interact with negatively charged nucleic acids under physiological conditions. They also have tertiary amines in the interior, which can promote the intracellular release of nucleic acids through the ‘proton sponge’ effect. However, dendrimer-based delivery of small interfering RNAs (siRNAs) into tumour stem cells is largely unexplored.

In this study, we demonstrate that knockdown of TLX using dendrimer nanovector-delivered synthetic siRNAs or virally expressed short hairpin RNAs (shRNAs) dramatically reduces GSC growth and self-renewal. By transplanting TLX shRNA-transduced GSCs into immunodeficient NOD SCID Gamma (NSG) mice, we show that knockdown of TLX leads to almost complete failure of GSCs to develop tumours in transplanted mouse brains. Furthermore, intratumoral delivery of TLX siRNAs using a dendrimer nanovector or TLX shRNAs using a viral vector inhibits GSC-induced tumorigenesis and prolongs the lifespan of GSC-grafted animals substantially.

Moreover, we identify TET3 as a critical TLX downstream target that inhibits GSC self-renewal and tumorigenesis.

Results
TLX shRNA reduces GSC self-renewal and tumorigenesis. To determine the role of TLX in human GSCs, we isolated ten primary GSC lines from tumour tissues of newly diagnosed human World Health Organization grade IV GBM patients and cultured them as three-dimensional tumour spheres in a culture condition for GSC enrichment. We classified these GSCs into GBM subtypes using a method previously reported. Among them, PBT003, PBT022, PBT726 and PBT1030 are classical, PBT017, PBT030 and PBT1008 are mesenchymal, whereas PBT024, PBT111 and PBT707 are proneural. These GSCs expressed human neural stem cell markers, Nestin and TLX (Supplementary Fig. 1a). They are also multipotent. When cultured in differentiation condition, they were able to differentiate into βIII tubulin-positive neurons and glial fibrillary acidic protein (GFAP)-positive astrocytes (Supplementary Fig. 1b).

To study the function of TLX in GSCs, two shRNAs were designed to knockdown TLX expression in GSCs with two scrambled RNAs as negative controls. After stably transducing TLX shRNA-expressing lentivirus into GSCs, efficient knockdown of TLX was confirmed (Fig. 1a). Knockdown of TLX expression dramatically reduced the growth rate of all GSC lines tested (Fig. 1b), indicating that TLX plays an important role in GSC expansion.

We next determined the effect of knocking down TLX on the self-renewal ability of GSCs using clonal analysis and limiting dilution assay. Knockdown of TLX dramatically reduced the self-renewal capacity of GSCs, as revealed by the sharply decreased sphere formation rate and stem cell frequency in TLX-shRNA-transduced cells (Fig. 2 and Supplementary Figs 2, 3). Together, these results indicate that TLX is essential for maintaining GSC growth rate and self-renewal ability.

The dramatic inhibitory effect of TLX shRNAs on GSC growth and self-renewal in vitro prompted us to test whether knockdown of TLX affects the ability of GSCs to form tumours in vivo. Two GSC lines, PBT003 and PBT707 cells, were transduced with a lentiviral vector expressing a TLX shRNA and a green fluorescent protein (GFP) reporter. The transduced cells were transplanted into the frontal lobe of NSG mouse brains. Tumour formation and expansion by the TLX shRNA-transduced GSCs were compared with scrambled control RNA-transduced GSCs. Although the same number of GFP-positive cells was injected into each mouse, GFP fluorescence imaging revealed large masses of GFP-positive cells in brains transplanted with control RNA-transduced GSCs, but there was barely any GFP signal in brains transplanted with TLX shRNA-transduced GSCs (Fig. 3a). Haematoxylin and eosin (H&E) staining showed that mice received control GSCs developed large tumour masses with typical infiltrative features of GBM (Fig. 3b, c). Conversely, GSCs treated with TLX shRNAs did not form tumours, or only formed small lesions that were confined to the injection sites (Fig. 3b, c). Stereological measurement of tumour volumes confirmed significantly smaller tumours in brains transplanted with GSCs treated with TLX shRNA, compared with that in brains transplanted with control GSCs (Fig. 3d).

Mice transplanted with TLX shRNA-transduced PBT003 cells had much better survival outcome compared with mice transplanted with scrambled control RNA-transduced cells (Fig. 3e). All mice that were transplanted with TLX shRNA-transduced PBT003 cells survived longer than mice transplanted with scrambled control RNA-transduced cells (Fig. 3e).
shRNA-transduced PBT003 cells survived for more than 70 days post transplant, and 40% of them survived beyond 80 days, whereas all mice that received PBT003 cells transduced with control RNA died before 60 days post transplant. Similarly, all mice that received PBT707 cells transduced with the control RNA died before day 85 post-transplant, whereas most mice that received PBT707 cells transduced with the TLX shRNA survived beyond this point, and 40% survived beyond day 110 (Fig. 3e). Together, these results indicate that knockdown of TLX suppresses tumour growth and increases the lifespan of GSC-grafted mice.

**Knockdown of TLX in vivo suppresses tumour progression.**

Next we tested if knocking down TLX in vivo could suppress the progression of human GSC-initiated tumours in a xenograft model. PBT003 cells were transduced with luciferase-expressing

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**Figure 1 | Knocking down TLX expression dramatically reduced the growth of GSCs.** (a) RT-PCR analysis of TLX expression in GSCs transduced with TLX shRNAs. Scrambled RNAs (SC1 and SC2) were included as negative controls. $N = 3$, *$P < 0.05$, **$P < 0.01$, ***$P < 0.001$ by Student’s t-test. Error bars are s.d. of the mean. (b) Growth kinetic analysis of the GSC lines transduced with scrambled control RNAs (SC1, SC2) or TLX shRNAs (shTLX1, shTLX2). $N = 4$. Error bars are s.d. of the mean.
lentivirus, which allowed us to monitor tumour growth in vivo by bioluminescence imaging. The resultant PBT003 cells were orthotopically transplanted into the frontal lobe of NSG mouse brains to establish tumours. One week after, mice were treated with scrambled control RNA or TLX shRNA-expressing lentivirus by intratumoral injection (Fig. 4a). Knockdown of TLX in PBT003 cells in vivo was confirmed by reverse transcription (RT)–PCR using human TLX-specific primers (Fig. 4b). Tumour formation was monitored using bioluminescence xenogen imaging (Fig. 4c). Mice receiving control RNA-expressing virus developed large tumours, whereas mice treated by TLX shRNA-expressing lentivirus had much smaller tumours (Fig. 4c,d). Bioluminescence measurement showed a significant decrease of tumour signal in mice treated with TLX shRNA-expressing virus at 5 weeks after treatment (Fig. 4d).

Moreover, PBT003-grafted mice treated with TLX shRNA-expressing virus had much better survival outcome compared with mice treated with scrambled control RNA (Fig. 4e). All mice that received control RNA died before day 60 post-treatment and the median survival was 56 days after viral treatment, whereas 60% of mice treated with TLX shRNA survived beyond 200 days post-treatment (Fig. 4e).

When mice in control group died, brain samples were collected for histological analysis. H&E staining revealed the development of big tumour mass and aggressive tumour invasion across the hemisphere in brains of control mice, whereas in brains of TLX

Figure 2 | Knocking down TLX expression dramatically reduced the self-renewal of GSCs. (a) Quantification of sphere formation rate of GSCs transduced with scrambled control RNAs (SC1, SC2) or TLX shRNAs (shTLX1, shTLX2). N = 6, **P < 0.01, ***P < 0.001 by Student’s t-test. Error bars are s.d. of the mean. (b) Limiting dilution assay (LDA) analysis of GSCs transduced with scrambled control RNAs (SC) or TLX shRNAs (shTLX). N = 20.
shRNA-treated mice collected at the same time, no or much smaller tumour was detected (Fig. 4f–h). The tumours developed in control mice exhibited typical infiltrative features of GBM (Fig. 4h). These results indicate that TLX shRNA-expressing virus suppressed the progression of established tumours and increased the lifespan of treated animals.

To determine the effect of TLX shRNA treatment on tumour progression at a different time point after tumour establishment, we treated mice with control RNA or TLX shRNA-expressing lentivirus 2 weeks after transplantation with luciferase reporter-bearing PBT003 cells (Supplementary Fig. 4a). Bioluminescence imaging showed that TLX shRNA-expressing lentivirus dramatically inhibited tumour growth, compared with control virus (Supplementary Fig. 4b). Bioluminescence measurement showed a significant decrease of tumour size in TLX shRNA-expressing lentivirus-transduced mice, at 3 and 5 weeks after treatment (Supplementary Fig. 4c). These results demonstrate that treatment with TLX shRNA-expressing lentivirus suppressed the progression of established tumours and increased the lifespan of treated mice. The above results together strongly support our hypothesis that TLX could be an effective target to suppress human GSC self-renewal and tumorigenesis.

The TLX siRNA nanocomplex inhibits GSC tumour progression.
In addition to knocking down TLX using a TLX shRNA-expressing viral vector, we explored delivering TLX siRNA oligonucleotides using a non-viral nanovector. We chose the poly(amidoamine) PAMAM dendrimer of generation 5 (referred to as G5 thereafter) to deliver TLX siRNA because it has been shown to deliver siRNAs effectively by forming stable and compact nanoparticles with siRNAs and protect siRNAs from degradation, leading to efficient and long-term gene silencing34,38. The dendrimer-mediated delivery is also relatively safe without discernible toxicity39.

We first tested whether the G5 dendrimer could form stable nanoparticles with siRNAs and deliver siRNAs into GSCs effectively. The G5 dendrimer could form stable complexes with TLX siRNA, as revealed by significant retardation on migration of the G5-siRNA complex at an N/P ratio of 1.0 or above in a gel shift assay (Supplementary Fig. 5a). G5 and TLX siRNA readily formed stable and uniform nanoparticles with an average size slightly smaller than 100 nm in diameter at N/P ratio of 5 (Supplementary Fig. 5b). Consistently, the G5-TLX siRNA complexes were able to protect siRNA from RNase-mediated degradation, whereas naked siRNA was rapidly degraded upon...
RNAse digestion (Supplementary Fig. 5c). When incubated with PBT003 cells, G5 efficiently delivered Cy3-labelled siRNA (Cy3-siRNA) into cells compared with Cy3-siRNA alone control (Fig. 5a). The cellular uptake of G5 delivered Cy3-siRNA was further confirmed by flow cytometry analysis (Fig. 5b).

To achieve tumour cell-targeted delivery, we coated the dendrimer-TLX siRNA nanoparticles with a targeting peptide that contains the dual targeting RGDK motif40. The RGDK motif directs tumour-specific homing through integrin-dependent binding to tumour cells specifically41, whereas the RXXK motif promotes cell and tissue penetration through interaction with neuropilin-1 (refs 41–44). Both integrin αv and neuropilin-1 were expressed on the surface of PBT003 cells (Fig. 5c and Supplementary Fig. 5d). Decoration of the TLX siRNA-G5 complexes with the RGDK peptide led to the formation of nanoparticles with a size ~100 nm in diameter (Supplementary Fig. 5e), within a size range required for effective cellular uptake. Coating the G5 dendrimer with the RGDK peptide enhanced the uptake of Cy3-siRNA into GSCs, compared with the uptake of G5 dendrimer-delivered Cy3-siRNA or Cy3-siRNA alone, as revealed by increased intensity of intracellular Cy3 fluorescence (Fig. 5d).

We then treated PBT003 cells with the G5 dendrimer-TLX siRNA nanocomplex with or without RGDK coating. RT–PCR confirmed efficient TLX knockdown by dendrimer-delivered TLX siRNA. Treatment with RGDK-coated dendrimer-TLX siRNA nanocomplex induced even more potent TLX knockdown, presumably due to better cell penetration (Fig. 5e). Compared with control RNA, TLX siRNA delivered by dendrimer dramatically reduced the growth of PBT003 cells, and TLX siRNA delivered by RGDK-coated dendrimer suppressed the growth of PBT003 cells even more (Fig. 5f). Together, these results demonstrated that RGDK-coated dendrimer-TLX siRNA complex efficiently knocked down TLX expression in GSCs and suppressed GSC growth potently.

Next we investigated whether TLX siRNA delivered by RGDK-coated dendrimer could suppress tumour progression in a human GSC-derived xenograft tumour model. PBT003 cells with a luciferase reporter were transplanted into the frontal lobe of NSG mouse brains to establish tumours. One week after transplantation, mice were treated with the RGDK-coated dendrimer-TLX siRNA or dendrimer-control RNA complex by intratumoral injection (Fig. 6a). The in vivo TLX knockdown was confirmed by RT–PCR (Fig. 6b). No obvious body weight loss was resulted from surgery or nanoparticle treatment before tumour-induced symptoms developed (Supplementary Fig. 6). Bioluminescence imaging revealed that mice treated with the RGDK-coated dendrimer-TLX siRNA complex had dramatically reduced tumour progression compared with control mice (Fig. 6c). Bioluminescence measurement confirmed that the tumour signals in siRNA complex-treated mice were significantly decreased compared with that in control mice (Fig. 6d). Moreover, treatment with the TLX siRNA complex significantly extended the lifespan of GSC-grafted mice (Fig. 6e). Tumours developed in control mice exhibited typical infiltrative features of GBM (Fig. 6f,g), whereas mice treated with the RGDK-dendrimer-TLX siRNA nanoparticles developed smaller tumour. These results indicate that RGDK-coated dendrimer-delivered TLX siRNA could effectively decrease tumour growth and increase the lifespan of tumour-bearing mice.

TET3 suppresses GSC self-renewal and tumorigenicity. To investigate the mechanism by which TLX controls the self-renewal and tumorigenesis of GSCs, microarray analysis was performed to compare gene expression profiles in control and TLX shRNA-treated GSCs. TLX was identified in the
downregulated gene cohort, whereas the cyclin-dependent kinase inhibitor p21, a known downstream target that is repressed by TLX\(^5\), was among the upregulated genes, confirming the effectiveness of TLX knockdown in TLX shRNA-treated cells. Several potential downstream targets that were not linked with TLX before were identified in our array analysis (Fig. 7a). Specifically, the expression of TET3, TDG and Dicer1 genes were upregulated and the expression of ID3, ID4 and MBD2 genes were downregulated in TLX-shRNA-treated PBT003 cells (Fig. 7a). The regulation of TET3 by TLX knockdown was confirmed in PBT003 cells and other GSC lines by RT–PCR (Fig. 7b,c and Supplementary Fig. 7a). Up-regulation of TET3 upon TLX knockdown was also confirmed in PBT003-grafted brain tumours from NSG mice treated with virus expressing TLX shRNA compared with that in tumours from control mice (Supplementary Fig. 8). Of note, TET3, TDG and MBD2 are all involved in DNA methylation modification, suggesting epigenetic regulation of DNA methylation may be an important downstream event of knocking down TLX in GSCs.

Based on the negative regulation of TET3 expression by TLX, we hypothesized that TET3 could function as a tumour suppressor to control GSC growth and self-renewal. To test this hypothesis, two shRNAs were designed to knockdown TET3 in GSCs. Knockdown of TET3 was confirmed in PBT003 and PBT707 cells (Supplementary Fig. 7b). PBT003 cells expressing TET3 shRNAs showed increased cell growth compared with control RNA-treated cells (Fig. 7d). Consistent with increased cell growth, PBT003 cells with TET3 knockdown also showed increased sphere formation rate compared with control cells (Fig. 7e). The increased cell growth and sphere formation rate after knockdown of TET3 were also observed in PBT707 cells (Fig. 7d,e). Taken together, these results demonstrated that knockdown of TET3 increased GSC growth and self-renewal.

To investigate if TET3 is sufficient to regulate the growth and self-renewal of GSC, we tested the effect of TET3 overexpression in GSCs. Human TET3 gene TET3-1 (with the CXXC domain) or TET3-2 (without the CXXC domain; Fig. 7f) was cloned into a lentiviral vector. Both TET3-1 and TET3-2 contain the dioxygenase domain that is present in all TET proteins. Overexpression of TET3-1 and TET3-2 was confirmed by RT–PCR in PBT003 and PBT707 cells transduced with the TET3-expressing virus (Fig. 7f and Supplementary Fig. 7c,d). Overexpression either TET3-1 or TET3-2 reduced the growth of both PBT003 and PBT707 cells (Fig. 7g). Consistent with decreased cell growth, PBT003 and PBT707 cells overexpressing TET3-1 or TET3-2 also showed decreased sphere formation rate compared with control cells (Fig. 7h). These results together indicate that TET3 inhibits GSC growth and self-renewal.

To investigate if TET3 regulates GSC tumorigenicity, PBT003 cells were transduced with a lentiviral vector expressing a TET3 shRNA or a control shRNA. The transduced cells were then transplanted into the frontal lobe of NSG mouse brains. Tumour formation and expansion by the TET3 shRNA-treated GSCs were compared with that by control RNA-treated GSCs maintained under identical conditions. Stereological measurement of tumour volumes confirmed the development of significantly larger

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**Figure 5 | Dendrimer-based nanoparticle delivery of siRNA into GSCs.**

(a) Cellular uptake of Cy3-siRNA delivered by siRNA alone or G5 dendrimers (at N/P ratio of 5) to PBT003 cells. Scale bar, 200 μm. Assays were repeated three times. (b) Cellular uptake of Cy3-siRNA analysed by flow cytometry. Uptake of Cy3-siRNA delivered by G5 dendrimer complexes into PBT003 cells was 98.4%. (c) Expression of integrin αv and neuropilin-1 on PBT003 GSCs revealed by immunostaining. Scale bar, 100 μm. (d) Cellular uptake of Cy3-labelled siRNA delivered by siRNA alone, G5 dendrimers or RGDK-coated G5 dendrimers (at N/P ratio of 5) to PBT003 cells. Scale bar, 200 μm. Assays were repeated three times. (e) Knockdown of TLX expression in PBT003 cells by the G5 dendrimer-TLX siRNA complex (G5-siTLX) and RGDK-coated G5-siTLX, analysed by RT-PCR. G5 dendrimer-scrambled control RNA (G5-SC) and RGDK-coated G5-SC nanoparticles were included as controls. (f) Growth inhibitory effect of the G5-siTLX and RGDK-G5-siTLX. PBT003 cells treated with G5 dendrimers or RGDK-coated G5 dendrimers complexes with SC or siTLX (at the N/P ratio of 5) were analysed by CellTiter-Glo luminescent assay. N = 3 for e; N = 6 for f. For both e and f, *P < 0.05, **P < 0.01, ***P < 0.001 by Student’s t-test. Error bars are s.d. of the mean.
Figure 6 | Treatment with the TLX siRNA nanoparticles reduced GSC-derived tumour growth and increased mouse survival. (a) Schematic of the experimental design, including GSC transplantation, dendrimer-siTLX complex treatment and xenogen imaging of xenografted tumours. The mice were treated with RGDK-coated G5 dendrimer-siTLX complex or RGDK-coated G5 dendrimer-SC complex. (b) In vivo knockdown of TLX using RGDK-G5-siTLX nanocomplex. \( N = 3 \), \(* * * P < 0.001\) by Student’s \( t \)-test. Error bars are s.e. of the mean. (c) Xenogen images of brain tumours in NSG mice treated with scrambled control (SC) or TLX siRNA (siTLX). The scale for bioluminescence intensity is shown on the right. (d) Quantification of the bioluminescence intensity of tumours treated with scrambled control (SC) or TLX siRNA (siTLX) in the brains of engrafted NSG mice. \( N = 7 \), \(* P < 0.05\), \(* * P < 0.01\) by Student’s \( t \)-test. Error bars are s.e. of the mean. (e) Survival curves of PBT003-engrafted NSG mice treated with scrambled control (SC) or TLX siRNA (siTLX). X axis represents days after dendrimer-siRNA treatment. \( N = 7 \) for each treatment group. \( P < 0.01\) by log-rank test. (f,g) H&E staining showing typical tumour infiltration characteristics of glioblastoma. Scale bar, 1 mm (f) and 50 \( \mu \)m (g). wk, week.

Tumours in brains transplanted with GSCs treated with TET3 shRNA, compared with that in brains transplanted with control GSCs (Fig. 7j). Mice transplanted with TET3 shRNA-transduced PBT003 cells had significantly shorter survival compared with mice transplanted with control RNA-transduced cells (Fig. 7k). Together, these results indicate that knockdown of TET3 increases tumour progression and decreases the lifespan of GSC-grafted mice, in a manner opposite to knockdown of TLX.
Figure 7 | TET3 regulates GSC self-renewal and tumorigenesis. (a) Microarray analysis of PBT003 cells transduced with virus expressing scrambled control (SC) or TLX shRNA (shTLX). (b, c) RT–PCR analysis showing TET3 upregulation upon TLX knockdown. N = 3, *P < 0.05, **P < 0.01 by Student’s t-test. Error bars are s.d. of the mean. (d, e) Cell growth (d) and sphere formation (e) analyses of the GSC lines transduced with control RNA (shC) or TET3 shRNAs (shTET3-1, shTET3-2). (f) Schematics of TET3-1 and TET3-2 proteins with characteristic domains. (g, h) Cell growth (g) and sphere formation (h) analyses of GSCs transduced with control (C) or TET3-expressing lentivirus, TET3-1 or TET3-2. (i) RT–PCR analysis showing overexpression of TET3 in GSCs. (j) Relative sizes (volumes) of brain tumours derived from PBT003 cells that were transduced with control RNA (shC) or TET3 shRNA (shTET3). (k) Survival curves of NSG mice transplanted with PBT003 cells transduced with control RNA (shC) or TET3 shRNA (shTET3). X axis represents days after GSC transplantation. For d, g, N = 4; for e, h, N = 6. Error bars are s.e. of the mean. *P < 0.05, **P < 0.01, ***P < 0.001 by Student’s t-test. For j, N = 3, ***P < 0.001 by Student’s t-test. Error bars are s.d. of the mean. For k, N = 8, P < 0.001 by log-rank test.
TET3 acts downstream of TLX to regulate GSC self-renewal.

To test whether TET3 acts downstream of TLX to regulate GSC growth and self-renewal, an inducible system to double knock down TLX and TET3 was established. PBT003 and PBT707 cells were transduced with lentivirus that expresses doxycycline (dox)-inducible TLX shRNA together with a puromycin-resistant reporter gene. After puromycin selection, the stably transduced cells were then transduced with lentivirus expressing dox-inducible TET3 shRNA. Induced knockdown of TLX and TET3 was confirmed in dox-treated PBT003 and PBT707 cells transduced with lentivirus expressing dox-inducible TLX shRNA (Fig. 8a) or TET3 shRNA (Fig. 8b). As expected, the expression of TET3 was upregulated after dox-induced knockdown of TLX (Fig. 8c). Dox-induced TET3 knockdown reversed the expression of TET3 to control levels in cells expressing both inducible TLX shRNA and inducible TET3 shRNA (Fig. 8c).

Next we tested whether inducible TET3 knockdown could rescue the inhibitory effect of inducible TLX knockdown on GSC growth and self-renewal. After dox induction, the growth of PBT003 and PBT707 cells expressing inducible TLX shRNA was reduced when compared with non-induced cells (Fig. 8d). The decreased cell growth resulted from induced TLX knockdown was rescued substantially by induced TET3 knockdown in both PBT003 and PBT707 cells (Fig. 8d). The reduced self-renewal of

![Figure 8](image-url)
PBT003 and PBT707 cells resulted from dox-induced TLX knockdown was also rescued by dox-induced TET3 knockdown (Fig. 8e). These results indicate that TET3 is a critical downstream target of TLX in regulating GSC growth and self-renewal.

The inverse correlation between TET3 and TLX expression in TLX knockdown GSCs led us to hypothesize that TLX, a known transcription factor that usually works as a transcriptional repressor \(^{45}\), could repress TET3 expression by directly binding to the promoter of the TET3 gene. Chromatin immunoprecipitation (ChiP) analysis using a TLX-specific antibody revealed that TLX bound to the promoter and proximal intron regions of TET3 containing the putative TLX-binding sites in both PBT003 and PBT707 cells (Supplementary Fig. 9a,b). At position 1 (P1, around the promoter region) and position 5 (P5, intron 2), the binding sites of TLX was associated with a higher level of repressive histone mark H3K9me3 than that of active histone mark H3K4me3 in both PBT003 and PBT707 cells (Supplementary Fig. 9c). These data suggest that TLX could regulate the transcription of TET3 by directly binding to regulatory regions of the TET3 gene.

To identify downstream targets of TET3 in GSCs, microarray analysis was performed. The gene expression profile of PBT003 cells treated with TET3 shRNA-expressing lentivirus was compared with PBT003 cells expressing control shRNA. Data sets from microarray analysis of PBT003 cells transduced with TLX shRNA or control RNA were included for comparison. Among the differentially expressed genes, an inverse correlation in gene expression was observed in TLX knockdown cells and TET3 knockdown cells (Fig. 8f), consistent with our hypothesis that TLX represses TET3 expression. Specifically, genes related to tumour-suppressive function, including BTG2, TUSC1, BAK1, LAT52, FZD6 and PPP2R1B, were upregulated in TLX knockdown cells, but downregulated in TET3 knockdown cells (Fig. 8g and Supplementary Fig. 10), suggesting that the TLX-TET3 regulatory cascade could regulate the growth and self-renewal of GSCs through regulating these downstream tumour suppressors.

Because TET3 is a dioxygenase that converts 5hmC to 5mC, we next tested whether the TLX-TET3 regulatory cascade could regulate 5hmC level in GSCs. Dot blot analysis using a 5hmC-specific antibody revealed increased 5hmC level upon TLX knockdown, but decreased 5mC level upon TET3 knockdown in PBT003 cells (Fig. 8h,i). Our DNA microarray analyses have identified several tumour suppressors as candidate downstream targets of TLX and TET3, including BTG2 and PPP2R1B (Fig. 8g), which were also upregulated upon TLX knockdown in PBT003-grafted brain tumours in NSG mice (Supplementary Fig. 8). To test whether the TLX-TET3 cascade regulates 5hmC level at the promoter of these potential downstream targets, hydroxymethylated DNA immunoprecipitation-quantitative PCR (qPCR) analysis was performed. The 5hmC levels at the promoter region of BTG2 and PPP2R1B genes were significantly increased in PBT003 cells transduced with TLX shRNA, compared with control cells (Fig. 8j,k), consistent with the elevated expression of these genes in TLX knockdown PBT003 cells (Supplementary Fig. 10a). In contrast, knockdown of TET3 reduced 5hmC level at the promoter region of BTG2 and PPP2R1B (Fig. 8j,k), consistent with the reduced expression of these genes in TET3 knockdown PBT003 cells (Supplementary Fig. 10b). These results indicate that the TLX-TET3 regulatory cascade could regulate the expression of downstream tumour-suppressor genes by controlling the 5hmC level at their promoter regions.

Discussion

Cancer stem cells are critical for tumour maintenance, metastasis and resistance to therapy. Therefore, targeting cancer stem cells is a priority in the development of novel cancer therapies that can completely cure and eradicate cancer by eliminating residual tumour-initiating cells. GSCs are implicated in the initiation and development of GBM, the most aggressive and invariably lethal brain tumour. This study identified the nuclear receptor TLX and TET3 regulatory axis as a target for GSCs. The results of this study demonstrated that TLX is fundamental for maintaining GSC growth, self-renewal and \textit{in vivo} tumour formation capacity, whereas TET3 is a potential tumour suppressor that inhibits GSC growth, self-renewal and tumorigenesis. Knockdown of TLX expression using either virally expressed shRNA or nanoparticle-delivered siRNA dramatically reduced the growth and self-renewal of GSCs \textit{in vitro} and impaired the ability of GSCs to form brain tumours \textit{in vivo}. The finding that \textit{in vivo} knockdown of TLX expression inhibits brain tumour development suggests that TLX is a promising target for anti-GBM therapy.

A role for TLX in glioblastoma development has been proposed in studies using mouse models \(^{14,46}\). A recent study by lineage tracing demonstrated that TLX regulates the self-renewal of brain tumour stem cells in mouse brains \(^{47}\). However, there is no study yet to directly investigate the role of TLX in human GSCs \textit{in vivo}. In this study, we showed direct evidence that targeting TLX in GSCs derived from human GBM patients efficiently inhibited the growth, self-renewal and tumorigenicity of GSCs \textit{in vitro} and \textit{in vivo}. The inhibitory effects of growth and self-renewal by TLX suppression were seen in all GSC lines tested, including classical, mesenchymal and proneural subtypes. These results suggest a general role for TLX in maintaining human GSC self-renewal, independent of GBM subtypes. Treatment of human GSC-grafted mice with TLX siRNA dramatically reduced tumour growth and significantly prolonged survival of GSC-grafted mice. Targeting TLX \textit{in vivo} is effective at different time points, both 1 week and 2 weeks after tumour establishment. Our study, by knocking down TLX \textit{in vivo}, provided proof-of-concept that targeting TLX is effective to suppress the progression of human GSC-derived tumours.

Our study, by knocking down TLX expression in GSCs, demonstrates that TLX is necessary for GSC self-renewal and tumorigenesis. Because TLX is an essential regulator of neural stem cell self-renewal \(^{11}\), it is possible that neural stem cells expressing high levels of TLX that beyond certain threshold may progress into gliomas over time. However, in a study using a transgenic model of TLX overexpression in neural stem cells, we did not detect any tumour formation \(^{48}\). In a separate study using a mouse model of TLX overexpression, no tumour formation was detected in brains of TLX-overexpressing mice under normal condition either \(^{14}\). Only when combined with loss of p53 or, to a less extent, ageing, overexpression of TLX led to glioma progression over time \(^{14}\). Therefore, although our study indicates that TLX is necessary for GSC self-renewal and tumorigenesis, overexpression of TLX alone may not be sufficient for GSC tumorigenesis.

Although the importance of targeting TLX in GBM is established, the knowledge on how TLX controls the self-renewal and tumorigenesis of human GSC is limited. The cyclin-dependent kinase inhibitor p21 is a known downstream target of TLX that can control cell cycle arrest. Our microarray confirmed that p21 is upregulated upon TLX knockdown as previously reported by us and others \(^{45,47}\). We did not see upregulation of reported tumour-suppressor genes, such as CDKN2A, CDKN2B and PML, and factors for neuronal differentiation, such as SMARCC1 and DLX2, in our microarray analysis. This may be due to the difference of human samples and mouse samples used for analyses. In our study, human GSCs after TLX knockdown were investigated, whereas previous study examined the expression of genes altered by TLX knockout.
using RNAs isolated from tumour tissues of a mouse tumour model. In this study, we identified TET3 as a potential tumour suppressor that acts downstream of TLX to regulate GSC growth and self-renewal. TLX represents the first transcription factor that has been identified to regulate TET3 expression. TET3 is a member of the TET family proteins that are known to be epigenetic regulators that control DNA demethylation. Although growing evidence showing that epigenetic regulation plays an important role in cancer development, our knowledge on the role of TET family members, especially TET3, in tumour development is rather limited. We show here that TET3 suppresses the growth, self-renewal and tumorigenesis of GSCs downstream of TLX. To our knowledge, this is the first study to define the role of TET3 in cancer stem cells and in GBM. Decreased 5mC level has been observed in human cancers, including malignant glioma, but the underlying mechanism remains to be discovered. Our finding showing that TLX represses TET3 expression in GSCs could provide a plausible explanation for the decreased 5mC level in GBM. Furthermore, we identified tumour suppressors, including BTG2, TUSC1, BAK1, LAT52, Fzd6 and PPP2R1B, as common targets of TLX and TET3. These tumour suppressors could be potential targets of the TLX-TET3 regulatory axis in GSCs that are worthy of further studies. The knowledge we gained about the role of TET3 in suppressing tumour stem cell growth and self-renewal, and the potential tumour-suppressor target genes we identified for TET3 in this study will trigger further studies to delineate the function of TET3 in tumour stem cells in particular and tumorigenesis in general.

Mammalian TET3 exists in isoforms either containing the CXXC domain (TET3-1) or not containing the CXXC domain (TET3-2) with, estimated transcript size of 11.6 and 10.9 kb, respectively. Of interest, the TET3-2 isoform that lack the CXXC amino-terminal CXXC domain, the TET3-2 form can also bind DNA. Although the TET3-1 isoform can bind to DNA using its amino-terminal CXXC domain, the TET3-2 form can also bind to DNA, presumably through its interacting proteins, including the CXXC domain containing protein CXXC4 (ref. 49), and transcription factors, such as REST. It is clear that the TET3-2 isoform that lacks the CXXC domain is able to induce 5mC formation and gene expression. Therefore, it is not surprising that the CXXC domain of TET3 seems dispensable for the effect on inhibition of GSC growth and self-renewal we observed in this study.

Small RNAs have gained increasing attention as candidate agents for therapies. However, the success of therapeutic application of small RNAs depends on efficient intracellular delivery. Safe and efficient small RNA delivery is in urgent need. In this study, we developed an efficient system of siRNA delivery into GSCs using polyacationic PAMAM dendrimer G5. This dendrimer has been shown to compact small RNAs into nanoparticles and protect RNAs from enzymatic degradation, therefore providing an efficient delivery means for introducing small RNAs into GSCs. Moreover, coating the dendrimer-siRNA nanoparticles with the tumour-homing RGDK peptide allows tumour-specific targeting. In this study, we show that GSCs express both integrin αv and neuropilin-1. Furthermore, we demonstrated that the RGDK-coated dendrimer-TLX siRNA nanoparticles were able to deliver TLX siRNAs into GSCs efficiently and exert potent gene knockdown and growth inhibitory effect. When delivered in vivo, the RGDK-coated dendrimer-TLX siRNA nanocomplex inhibited the growth of human GSC-initiated tumours. Moreover, treatment of the RGDK-coated G5-TLX siRNA complex significantly extended the lifespan of tumour-bearing mice, increasing the median survival from 48 days after the first treatment to 53 days, corresponding to about 7-month prolonged survival in patients. Similarly, mild but significant increase of survival has also been observed by targeting other important GBM targets. A potential clinical significance of this finding is derived from the ability of knocking down TLX in GSCs via lentivirus-delivered shRNA or nanoparticle-delivered siRNA to compromise the self-renewal and tumour formation potential of GSCs in vivo. Our studies demonstrated that TLX knockdown inhibited tumour initiation and progression from human GSCs in a xenografted tumour model and increased the survival of grafted animals substantially. GBM is highly aggressive brain tumour with a short life expectancy of a little over a year after diagnosis, and patient survival is only marginally increased by current therapies. The small RNA approach to knock down TLX in GSCs has the potential to help improve the outcome and survival of GBM patients.

Methods

Cell culture. Sphere cultures of GSCs were established from freshly dissociated spherical specimens as described. Patients were newly diagnosed as grade IV GBM multiforme based on World Health Organization-established guidelines. All patient tissues were obtained in accordance with the City of Hope Institutional Review Board-approved protocols. The study uses completely anonymized specimens. No informed consent is involved. The GSCs were maintained in DMEM-F12 medium (Omega Scientific) supplemented with 1X B27 (Invitrogen), 5 μg ml−1 heparin (Sigma), 2 mM l-glutamine (Media Tech), 27.4 μM HEPEs (Fisher), 20 ng ml−1 EGF (PeproTech) and 20 ng ml−1 FGF (PeproTech) with growth factors replenished twice a week. GSCs were treated with actussce (Innovative Cell Technologies) for cell dissociation and induced into differentiation using 0.5% fetal bovine serum and 1 μl all-trans retinoic acid. All the cultures used in this study were confirmed for the lack of mycoplasma contamination using MycoAlert PLUS Mycoplasma Detection Kit (Lonza).

Plasmid DNA and viral preparation and transduction. shRNAs or the scrambled plasmid RNAs were cloned into lentiviral pHIV7-GFP or pHIV-TetR-GFP vector. The sequences for shRNAs include TLX shRNA-1 (5'-GGGCGCATGCTGCGGCGAGT-3') and TLX shRNA-1 scrambled control (5'-CAGTGGCATGACGCCTGGCGT-3'), TLX shRNA-2 (5'-GGAAATTGACATGACAAAAGA-3') and TLX scrambled shRNA-2 (5'-ACTCCAAAGGAGTCAAGAGA-3'), shRNA control for TET3 (5'-GTTCCAGATGTTGGGCGAGT-3') shTET3-1 (5'-CCGAAAGCTGTGTCCTTCTTA-3') and shTET3-2 (5'-GGGATCCACTCTTCTAGTAC-3'). Lentiviruses were produced using 293T cells. To transduce GSCs, cells were dissociated and reseeded with lentiviruses and 4 μg ml−1 polybrene for 24h. Human TET3-expressing vector EX-H2292-M1 was purchased from GeneCopoeia and HEK293T sequences were subcloned into CSC lentiviral vector to get TET3-2-expressing vector. Human TET3 CXXC domain sequences were amplified from human neural stem cell cDNAs and subcloned into TET3-2 expressing vector to get TET3-1 expressing vector that expresses full length of TET3 gene.

RT-PCR. Total RNAs were isolated with Trizol reagent (Invitrogen) or RNeasy Mini Kit (Qiagen). Reverse transcription (RT) was performed using the Tetro cDNA synthesis Kit (BioLINE). RT–PCR reactions were performed using SYBR Green Master Mix (Thermo Scientific) on Step One Plus Real-Time PCR instrument (Applied Biosystems). The primers for RT–PCR are listed in Supplementary Table 1. Actin or glyceraldehyde-3-phosphate dehydrogenase was used as the reference gene for normalization. The ΔΔCt method was used for quantification analysis.

Immunostaining. For immunofluorescence, cells were fixed with 4% paraformaldehyde and permeabilized in 0.3% Triton X-100. Antibodies included rabbit anti-TLX (1:1,000; Shl lab)52, mouse anti-nestin (1:2,000; BD Pharmingen; Catalogue # 611659)52, rabbit anti-integrin αv (1:500; Chemicon; Catalogue # AB192513), mouse anti-neuropilin-1 (1:200; Milleniy Biotech; Catalogue # 130-090-693)52, mouse anti-VEGF (1:1,000; Sigma; Catalogue # G3893) and rabbit anti-Tuj1 (1:6,000; Covance; Catalogue # PRB-435P)52.

Animals. All animal-related work was performed under the IACUC protocol approved by the City of Hope Institutional Animal Care and Use Committee. Both male and female NSG mice (the Jackson Laboratory) at 6–8 weeks old were used at age- and gender-matched manner. The sample size was determined based on using t-test for two-group independent samples to reach power of 0.8 and the significance level of 0.05. P < 0.05 was considered statistically significant. When monitoring tumour growth, investigators were blind to the group
allocation during the bioluminescence xenogen imaging and aware of group allocation when assessing the outcome.

**Viral transduction followed by transplantation.** GSCs were transduced with control RNA or relevant siRNA expressing lentiviruses. Two days after virus transduction, $5 \times 10^5$ cells were transplanted into the frontal lobes of brains of 6- to 8-week-old NSG mice by stereotaxic intracranial injection. Briefly, 2 μl dissociated cells in PBS were injected into the following site (AP = 0.6 mm, ML = ±1.6 mm and DV = −2.6 mm) with a rate of 1 μl/min. The same coordinates were used for all viral intracranial injections in this study. Mouse brains were harvested when severely sick mouse was found in treated or control group. H&E staining was performed on 20 μm coronal sections of frozen brain samples, followed by tumour size analysis. The tumour volume was measured by multiplying the area of the tumour tissues (quantified by Image J) by the thickness of the sections, then multiplying by the number of the sections that contain the tumour tissues. In a separate set of experiment, the survival of grafted mice was recorded and analysed.

**Intracranial viral transduction.** PBT003 cells ($2 \times 10^5$) transduced with luciferase-expressing lentivirus were intracranially transplanted into the frontal lobe of 6- to 8-week-old NSG mice. One week or two weeks later, mice were randomly grouped and treated with scrambled control RNA or TLX siRNA-expressing lentivirus by intratumoral injection. Tumour growth was monitored by bioluminescence xenogen imaging and aware of group allocation during the bioluminescence xenogen imaging and aware of group allocation when assessing the outcome. The same coordinates were used for all viral intracranial injections in this study. Mouse brains were harvested when severely sick mouse was found in treated or control group. H&E staining was performed on 20 μm coronal sections of frozen brain samples, followed by tumour size analysis. The tumour volume was measured by multiplying the area of the tumour tissues (quantified by Image J) by the thickness of the sections, then multiplying by the number of the sections that contain the tumour tissues. In a separate set of experiment, the survival of grafted mice was recorded and analysed.

**Dendrimer-based siRNA delivery in vitro.** The G5 dendrimer was synthesized starting with the triethanolamine core and following the iterative Michael addition and amidiﬁcation as previously described. The RGDK peptide was synthesized as an oligopeptide with the sequence of E16G6RGDK40 and purchased from GL Biochem Ltd. The TLX siRNA has the sense of sequence: 5'-CCGCAAUUUGACGGCCUUCAGAUAUdgA-3', antisense: 5'-UCAUCAUCUUGAAGGGCGUCAUUAUGGGGGG-3'. The control siRNA has the sense of sequence: 5'-CAUCAUCAUCUUGAAGGGCGUCAUUAUGGGGGG-3', antisense: 5'-UCAUCAUCUUGAAGGGCGUCAUUAUGGGGGG-3'. To form RGDK-coated G5 dendrimer-siRNA complexes, G5 was first mixed with siRNAs at N/P (total terminal amines in G5)/[phosphates in siRNA]) ratio of 5 and kept at 37°C for 30 min. Then RGDK peptide was added to the G5-siRNA complex at G5/RGDK molar ratio of 0.426 and incubated at 37°C for another 10 min. The G5-siRNA or RGDK-G5-siRNA complexes were then added to GSCs. PBT003 cells ($2 \times 10^5$) were treated with Cy3-labelled siRNA delivered by siRNA alone, G5 dendrimers or RGDK-coated G5 dendrimer-siRNA (N/P ratio of 5; G5/RGDK ratio of 0.426). Two days after, cellular uptake of Cy3-labelled siRNA delivered by siRNA alone, G5 dendrimers or RGDK-coated G5 dendrimers was monitored by fluorescence microscopy and flow cytometry. For knocking down of TLX, PBT003 cells were treated with the G5 dendrimer-TLX siRNA complex or G5-control siRNA complex with or without RGDK coating, TLX expression was analysed by RT–PCR after 2 days of dendrimer-TLX siRNA treatment.

**Intracranial delivery of dendrimer-siRNA nanoparticles.** PBT003 cells ($2 \times 10^5$) transduced with luciferase-expressing lentivirus were intracranially transplanted into the frontal lobe of 6- to 8-week-old NSG mice. One week after transplant, tumours were detected by bioluminescence imaging and mice were treated with RGDK-coated G5 dendrimer-TLX siRNA complex or RGDK-coated G5 dendrimer-SC complex (2.5 nmole siRNA per mouse with N/P ratio of 5; G5/RGDK ratio of 0.426) by intratumoral injection once a week for 6 weeks. Tumour growth was monitored by bioluminescence imaging once a week for 7 weeks. The bioluminescence intensity, mouse body weight and mouse survival were analysed.

**Microarray analysis and GBM subtype determination.** For GBM subtype characterization, total RNAs were extracted from ten lines of GSCs using RNeasy Mini Kit (Qiagen). Microarray analysis was performed using GeneChip Human Genome U133A 2.0 Array (Affymetrix). Microarray labelling, hybridization and quality control measurements were performed in the Integrative Genomics Core of City of Hope. The microarray expression data of the ten GSC samples were pooled together with published microarray expression data of 173 TCGA samples. After batch removal, principle component analysis (PCA) was performed on the pooled data set using Partek Genomics Suite software, version 6.6 (2014 Partek Inc). 747 relevant genes were used for PCA according to the data filtering approach as described. Centroids of the four GBM subtype clusters were defined by PCA and subjected to dot blot analysis using an antibody specific for 5hmC (1:5,000; Active Motif; Catalogue # 39770) for each reaction. The immunoprecipitation was carried out using magnetic beads conjugated protein G (Thermo Fisher). Primers used are listed in Supplementary Table 1.

**3hmC dot blot.** PBT003 cells were transduced with relevant siRNA-expressing lentivirus. Three days after, genomic DNA was extracted (QiAamp DNA Mini Kit) and subjected to dot blot analysis using an antibody specific for 5hmC (1:5,000; Active Motif; Catalogue # 39770) for each reaction. The immunoprecipitation was carried out using magnetic beads conjugated protein G (Thermo Fisher). Primers used for BTG and PPP2R1B RT–PCR are listed in Supplementary Table 1.
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Author contributions
Y.S. conceived the project; Q.C., S.Y., P.Y., E.T., Guo.S., J.Z., Gui. S., X.L., C.C., K.M., C.Z., K.T.A., M.D., C.B., B.B. and Y.S. performed experiments; L.Y., C.W. and X.W. analysed microarray data; Q.C., S.Y., L.P., A.R., J.J.R. and Y.S. designed experiments and interpreted results; Q.C. and Y.S. wrote the manuscript with comments from all other authors.

Additional information
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