Efficient gene silencing in brain tumors with hydrophobically modified siRNAs

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**Conflict of Interest**

A.K. discloses ownership of the stock of RXi Pharmaceuticals and Advirna, LLC. Other authors have no conflict of interest to report.

**Abstract**

Glioblastoma (GBM) is the most common and lethal form of primary brain tumor with dismal median and two-year survivals of 14.5 months and 18%, respectively. The paucity of new therapeutic agents stems from the complex biology of a highly adaptable tumor that uses
multiple survival and proliferation mechanisms to circumvent current treatment approaches. Here, we investigated the potency of a new generation of small interfering RNAs (siRNAs) to silence gene expression in orthotopic brain tumors generated by transplantation of human glioma stem-like cells (GSCs) in athymic nude mice. We demonstrate that cholesterol-conjugated, nuclease-resistant siRNAs (Chol-hsiRNAs) decrease mRNA and silence luciferase expression by 90% in vitro in GBM neurospheres. Furthermore, Chol-hsiRNAs distribute broadly in brain tumors after a single intratumoral injection, achieving sustained and potent (>45% mRNA and >90% protein) tumor-specific gene silencing. This readily available platform is sequence-independent and can be adapted to target one or more candidate GBM driver genes, providing a straightforward means of modulating GBM biology in vivo.

**Introduction**

Glioblastoma (GBM, or grade IV astrocytoma) is the most frequent and lethal primary malignant brain tumor in humans, characterized by invasiveness, angiogenesis, and frequent necrosis(1). Current treatments are limited to surgical tumor resection, followed by chemotherapy with temozolomide and radiotherapy, and do not greatly improve prognosis(2,3). GBM recurrence is almost inevitable, due in part to stem-like tumor cells (GSCs, or tumor-initiating cells) that are both highly invasive and resistant to chemotherapy and radiation(4). These challenges underscore the need for novel approaches to GBM treatment.

The massive repository of information available from cancer genomics studies and the ever-expanding knowledge about the role of glioma stem-like cells (GSCs) in tumor biology have revealed a large number of genes and networks that orchestrate many of the pernicious properties of GBM(5-7). However, small molecule drugs, which have been the mainstay of the
pharmaceutical industry, can only target 10-15% of proteins encoded in the genome, and primarily those with enzymatic activity(8). A considerable number of genes (e.g. transcription factors, histones, extracellular matrix proteins), long non-coding RNAs (lncRNAs), and microRNAs (miRNAs) critical to GBM biology are not targetable by this approach. In contrast, siRNAs and antisense oligonucleotides (ASOs) can target any coding (mRNA) or non-coding RNA (lncRNA, miRNA)(9). Therapeutic oligonucleotides are a highly flexible platform because target specificity is defined by the base sequence and can to some extent be optimized independently from the PK/PD properties (defined by the backbone chemistry and targeting ligand); this separation does not exist for small molecule drugs, where the structure determines both properties(9). Therefore, oligonucleotide therapeutics have the necessary combination of specificity and flexibility to develop potent network (multi-mechanistic) drugs to effectively control GBM tumors(10).

Here, we investigated the activity of a next-generation siRNA (Chol-hsiRNAs)(11) in GBM8 cells, a patient-derived tumor-initiating GSC line(12). Chol-hsiRNAs are asymmetric siRNAs with a short duplex region (15 base pairs) and a single-stranded, fully phosphorothioate-modified tail(13). The 2ʹ-hydroxyl group in each nucleotide is substituted with 2ʹ-fluoro or 2ʹ-O-methyl modifications in an alternating pattern to provide stability and block innate immune activation(11,14-16). The 3ʹ end of the passenger strand is conjugated to cholesterol to enhance cellular uptake(11). Previously, we have shown that Chol-hsiRNAs are rapidly internalized by neural cells both in vitro and in vivo and induce potent gene silencing in a carrier-free manner(11). Here, we show that Chol-hsiRNAs are also rapidly internalized into GBM8 cells and induce potent mRNA and protein silencing both in vitro and in vivo in established orthotopic brain tumors. Chol-hsiRNAs are informational drugs that can be used to target any sequence and
thus can effectively manipulate gene expression in GBM, supporting the development of powerful new research tools and therapeutics for these devastating tumors.

**Materials and Methods**

**Oligonucleotide Synthesis**

Oligonucleotides were synthesized on an Expedite ABI DNA/RNA Synthesizer following standard protocols. Each synthesis was done on a 1 µmole scale using cholesterol-conjugated CPG for the sense strand and Unylinker solid support (ChemGenes, Wilmington, MA) for the antisense strand. All phosphoramidites (2´-O-methyl (ChemGenes, Wilmington, MA), 2´-fluoro (BioAutomation, Irving, Texas), and Cy3 (Quasar570) (Gene Pharma, Shanghai, China)) were prepared as 0.15 M in acetonitrile (ACN). Phosphoramidite coupling time was 4 min using 30 equiv. of the monomer. 5-(Ethylthio)-1H-tetrazole (ETT) 0.25 M in ACN was used as coupling activator. Detritylations were performed using 3% dichloroacetic acid (DCA) in dichloromethane (DCM) for 80 s, capping was done with acetic anhydride/THF/2,6-Lutidine, (80/10/10, v/v/v) (CAP A), and 1-methylimidazole/THF (16/84, v/v) (CAP B) for 15 s, and oxidation was achieved with 0.02 M iodine in THF/pyridine/water (70/20/10, v/v/v) for 80 s. Phosphorothioate linkages were introduced using 0.1 M solution of DDTT in ACN for 3 min.

**Oligonucleotide Deprotection and Purification**

Both sense and antisense strands were cleaved and deprotected using 1 mL of 40% aq. methylamine at 65°C for 15 min. The crude oligonucleotides were cooled, frozen in dry ice, and dried in a speedvac overnight. The resulting pellets were resuspended in 1 mL of 5% ACN. Antisense strand purification was performed on an Agilent 1100 series system equipped with an
Agilent PL-SAX, a polymer ion exchange column (4.6 x 150 mm), using the following conditions: Eluent A: 30% ACN and Eluent B: 1M sodium perchlorate in 30% ACN, gradient: 0% B for 2 min, 0-10% B for 1 min, 35% B for 8 min, equilibration to initial conditions for 6 min. Sense strand purification was performed on an Agilent 1100 series system equipped with a PRP-C18, a polymer reverse phase column (4.6 x 150 mm), using the following conditions: Eluent A: 50 mM sodium acetate in 5% ACN and Eluent B: ACN, gradient: 0% B for 2 min, 0-40% B for 1 min, 40-70% B for 8 min, equilibration to initial conditions for 6 min. Temperature was held at 70°C and flow rate at 10 ml/min for both cases. Peaks were monitored at 260 nm. Purified oligonucleotides were collected, frozen in dry ice, and dried in a speedvac overnight. Oligonucleotides were re-suspended in 5% ACN, desalted through fine Sephadex® G-25 columns built in house, lyophilized, and stored at -80°C until further use.

**LC-MS Analysis of Oligonucleotides**

The identity of oligonucleotides were established by LC-MS analysis on an Agilent 6530 accurate mass Q-TOF LC/MS machine using the following conditions: Buffer A: 100 mM HFIP/9 mM TEA in LC/MS grade water, Buffer B: 100 mM HFIP/9 mM TEA in LC/MS grade methanol, column: Agilent AdvanceBio oligonucleotides C18, gradient antisense: 0% B for 1 min, 0-30% B for 8 min, equilibration for 4 min, gradient sense: 0% B for 1 min, 0-50% B for 0.5 min, 50-100% B for 8 min, equilibration for 4 min, temperature: 45°C, flow rate: 0.5 mL/min, UV (260nm). MS parameters, Source: ESI, ion polarity: negative mode, range: 100-3200 m/z, scan rate: 2 spectra/s, VCap: 4000, fragmentor: 180V.
Cell Culture

GBM8 primary human glioblastoma cells were received as frozen stocks in 2014 from Dr. Miguel Sena-Esteves (University of Massachusetts Medical School), who originally obtained them from Dr. Samuel Rabkin (Massachusetts General Hospital, Boston, MA)(12). Cells were certified mycoplasma negative by regular testing (MycoAlert, Lonza), with the last date of testing conducted two weeks prior to the first in vitro experiment. Cell authentication was not performed prior to conducting the experiments described in this paper. The GBM8-fLuc cells used in these studies(17) were grown in suspension in Neurobasal media (Gibco) supplemented with 3 mM L-Glutamine (CellGro), 1X B27 supplement (Gibco), 0.5X N2 supplement (Gibco), 2 μg/ml heparin (Sigma), 0.5X antibiotic-antimycotic solution (CellGro), 0.5X amphotericine B (CellGro), 20 ng/ml recombinant human basic fibroblast growth factor (Peprotech), and 20 ng/ml recombinant human epidermal growth factor (Peprotech). Cell cultures were fed with 1/3 volume of fresh medium every other day and passaged once per week via neurosphere dissociation. Cell cultures were maintained for 1-2 passages (roughly two weeks) after thawing prior to use in all in vitro and in vivo experiments.

Oligonucleotide Delivery to GBM Neurospheres In Vitro

For efficacy studies, GBM8-fLuc cells were plated at 50,000 cells per well in 96-well tissue culture plates. Chol-hsiRNAs were diluted to twice the final concentration in OptiMEM (Gibco), and 50 μL diluted Chol-hsiRNA was added to 50 μL of cells, and were subsequently incubated for 72 h at 37°C and 5% CO₂. mRNA levels were quantified from cells using the QuantiGene 2.0 DNA Assay (Affymetrix) as described previously(18). Luciferase levels were quantified using a modified Dual-Glo Luciferase Assay System (Promega). Briefly, 50 μL of Dual-Glo Luciferase
Assay Reagent was added to each well, incubated for 10 minutes, and luminescence was recorded on an Infinite M1000 Pro microplate reader (Tecan). For fluorescence imaging studies, GBM8-fLuc cells were plated at 100,000 cells/mL in 6-well tissue culture plates. Cells were counterstained with NucBlue Live Cell Stain ReadyProbes reagent (Life Technologies) according to the manufacturer’s recommended protocol. Cy3-labeled Chol-hsiRNAs were added to the wells to a final concentration of 0 – 0.75 μM and incubated for 0 – 5 h at 37°C and 5% CO₂. Neurospheres were collected by centrifugation, affixed them to slides using a Shandon EZ Double Cytofunnel, fixed in 4% paraformaldehyde, and imaged on a Leica DMi8 inverted fluorescence microscope (40X oil objective). Intensity and exposure settings were the same for all samples. Images were processed using ImageJ (1.47v) software.

**Tumor Engraftment in Mouse Brain**

Athymic nude female mice (6-8 weeks old) were used in all studies. The humane endpoint for xenografted mice was defined as the loss of >15% of maximum body weight or appearance of any moribund symptoms (such as hunched posture). All animal procedures were approved by the University of Massachusetts Medical School Institutional Animal Care and Use Committee (IACUC, protocol number A-2061).

On the day of surgery, GBM8-fLuc cells were dissociated and prepared as a single cell suspension in sterile Dulbecco’s phosphate-buffered saline (PBS, Gibco). Animals were anesthetized and 50,000 cells in 1 μL were injected by stereotactic placement into the right striatum (coordinates relative to bregma: +0.5 mm AP, -2.0 mm ML, and -2.5 mm DV from skull surface) at an infusion rate of 0.125 μL/min. The kinetics of tumor growth were monitored by *in
in vivo imaging of tumor associated bioluminescence signal using an IVIS 100 (Perkin-Elmer) as described (TABS, Table S1). (17) Data acquisition and quantification were performed using Living Image 4.2 software (Perkin-Elmer).

**Oligonucleotide Delivery to Established Xenografts in Mouse Brain**

Tumor growth was monitored by *in vivo* bioluminescence imaging of tumor-associated luciferase activity (TABS) as described, and tumor-bearing mice assigned to groups with identical average TABS(17). Tumor-bearing athymic nude female mice (6-8 weeks old) were anesthetized and received stereotaxic injections of Chol-hsiRNAs into either the tumor (same coordinates used for tumor implantation) or the ipsilateral ventricle (coordinates relative to bregma: -0.2 mm AP, -0.8 mm ML, and -2.0 mm DV from skull surface).

For biodistribution studies, mice were injected intratumorally as described previously. Animals with roughly equivalent TABS (500,000 and 461,000 p/sec/cm²/sr) were selected to evaluate the distribution of the fluorescently labeled Chol-hsiRNA within the tumor microenvironment (Table S1). After 48 hours, mice were euthanized and perfused with PBS and 4% paraformaldehyde solution, and brains were post-fixed in 2% paraformaldehyde for 48 hours. Paraffin 4 μm coronal brain sections through the striatum were stained with hematoxylin and eosin for neuropathological analysis and with DAPI for biodistribution analysis of Cy3-labeled Chol-hsiRNAs in tumors. Images were acquired on a Leica DMi8 inverted microscope (40X).

For intratumoral efficacy studies, animals (n = 5 per group) were injected with 2 μl of artificial cerebrospinal fluid (aCSF – 120 mM NaCl, 25 mM NaHCO₃, 2.5 mM KCl, 1 mM NaH₂PO₄, 1.3
mM MgCl$_2$, 10 mM glucose), Chol-hsiRNA$^{fLuc}$ (31-62 μg, 2.5-5 nmol), Chol-hsiRNA$^{HTT}$ (62 μg, 5 nmol), or Chol-hsiRNA$^{NTC}$ (62 μg, 5 nmol) at an infusion rate of 0.125 μl/min. For intracerebroventricular efficacy studies, animals (n = 5 per group) were injected with 5 μl of aCSF, Chol-hsiRNA$^{fLuc}$ (31-62 μg, 2.5-5 nmol), or Chol-hsiRNA$^{NTC}$ (62 μg, 5 nmol) at an infusion rate of 0.5 μl/min.

**Oligonucleotide-Mediated Silencing of mRNA and Protein In Vivo**

For efficacy studies, brains were collected one week post-injection. Three 300 μm coronal sections through the striatum were collected and from each section a 2 mm punch was taken from the ipsilateral striatum and placed in RNAlater (Ambion) for 24 hours at 4°C. For mRNA quantification, each punch was lysed and processed as an individual sample for Quantigene 2.0 assay analysis (Affymetrix), and averaged for a single animal point(18). Luciferase activity in tissues was measured using the Promega Dual-Glo® Luciferase Assay System, as described earlier(19), and normalized to total protein content measured by a Bradford assay.

**Statistical analysis**

Data were analyzed using GraphPad Prism 6 software (GraphPad Software, Inc., San Diego, CA). For each independent mouse experiment, the level of mRNA silencing at each dose was normalized to the mean of the aCSF-injected control group. In vivo data were analyzed using a one-way ANOVA with a post hoc Bonferroni multiple comparisons test.

**Results**
Human primary glioma neurospheres productively internalize cholesterol-conjugated hsiRNA (Chol-hsiRNAs) in vitro

We evaluated the efficacy of Chol-hsiRNA (Fig. 1a) to silence gene expression in GBM. For these studies, we used GBM8, a well-validated GBM-initiating stem-like cell line that recapitulates the migratory and invasive properties of human glioblastoma following implantation into the brain of immunocompromised mice(12,20,21). First, we assessed the activity of Chol-hsiRNAs towards GBM8 neurospheres in vitro (Table S2). When added directly to the media, Chol-hsiRNAs were rapidly internalized by GBM8 neurospheres, over a period of hours (Fig. 1b). Chol-hsiRNA uptake is productive, as treatment of cells with hsiRNAs targeting PPIB mRNA (Cyclophilin B; PPIB-567) or HTT mRNA (Huntingtin; HTT-10150) for 72 hours reduced target mRNA levels by more than 90% (Fig. 1c) and 65%, respectively (Fig. 1d). No off-target silencing of either gene was observed following treatment with a non-targeting (scrambled) Chol-hsiRNA control (Chol-hsiRNANTC, Fig. 1c, 1d, gray bar, 3 µM). This cell line was transduced with the lentivirus vector CSCW2-fLuc-IRES-mCherry to generate GBM8-fLuc cells that constitutively express firefly luciferase and mCherry protein(22). Using a fLuc-targeting Chol-hsiRNA (Chol-hsiRNAfLuc), we assessed silencing of luciferase mRNA and protein at 72 hours after treatment. We observed an 82% reduction in luciferase mRNA (Fig. 2a), and a concomitant 90% reduction in luciferase activity (Fig. 2b). To ensure these effects were not due to compound toxicity, we assessed luciferase activity over a broad range of Chol-hsiRNA NT C concentrations (0.1 µM – 6 µM), and found no significant decrease in luciferase activity compared to the untreated control (Fig. 2c). We did not observe any significant reduction in GBM8 cell viability following administration of up to 5 µM Chol-hsiRNA (Fig. S1, Supplementary Methods). Additionally, we monitored neurosphere morphology at the highest
doses of each compound (Fig. 2d, 3 µM Chol-hsiRNA<sup>Luc</sup>; 6 µM Chol-hsiRNA<sup>NTC</sup>). There were no striking visual differences in neurosphere size (~250 µm) or number, suggesting that Chol-hsiRNA treatment did not trigger GBM8 cell death. Thus, Chol-hsiRNAs provide a simple and straightforward approach to achieve potent and specific mRNA silencing in GBM8 cells.

**Chol-hsiRNA distributes throughout brain tumors after direct injection**

We next interrogated the ability of Chol-hsiRNAs to distribute and silence gene expression in orthotopic GBM8 tumors in mouse brain. Two weeks after GBM8-fLuc tumor cell implantation, we injected 5 nmol Cy3-labeled Chol-hsiRNA<sup>HTT</sup> (5 nmol, 62.5 µg) into the tumor and analyzed distribution after 48 hours. As previously reported(17), GBM8-fLuc cells infiltrated the right hemisphere and exhibited characteristic hypercellular morphology with hyperchromatic nuclei, frequent mitoses, and necrosis (Fig. 3a). Cy3-labeled Chol-hsiRNA<sup>HTT</sup> distributed broadly throughout the main tumor mass (Fig. 3b, dotted white line). At higher magnification, we observed Cy3-labeled Chol-hsiRNA<sup>HTT</sup> accumulation in cytoplasmic foci, in addition to strong staining of the extracellular matrix and axonal bundles (Fig. 3b, arrows). This distribution pattern has been observed previously for Chol-hsiRNA following intraparenchymal injection into wild-type mouse striatum(11). We concluded that Chol-hsiRNAs distribute throughout established GBM8 tumors and are effectively taken up by tumor cells following a single, intratumoral injection.

**Intratumoral injection of Chol-hsiRNA potently silences gene expression in brain tumors**

Next, we analyzed the ability of Chol-hsiRNAs to silence gene expression in GBM8 tumors at two and four weeks after implantation. In these studies, we targeted two genes
selectively expressed in the tumor (human \( HTT \) and \( fLuc \)), to distinguish from silencing in normal mouse brain cells. We first sought to determine whether intratumoral or intracerebroventricular injection was a superior route of administration to silence gene expression in established brain tumors. Tumor-bearing mice received either an intratumoral or intracerebroventricular injection of Chol-hsiRNA\(^{HTT}\) four weeks after tumor implantation. One week after injection, we observed a 45% reduction in human \( HTT \) following a 2 nmol intratumoral injection and a 23% reduction in human \( HTT \) following a 5 nmol ICV injection, compared to a control injected with artificial cerebral spinal fluid (aCSF) (Fig. 4a). As a single intratumoral injection achieved greater levels of mRNA silencing, we proceeded with that route of administration for the subsequent studies.

Next, we assessed protein silencing after intratumoral delivery of Chol-hsiRNA\(^{fLuc}\) to two-week old GBM8-fLuc tumors. One week after intratumoral injection of either 2.5 or 5 nmol of Chol-hsiRNA\(^{fLuc}\), we observed a ~90% reduction in luciferase activity relative to the aCSF-injected control (Fig. 4b). The heterogeneity in luciferase activity for the Chol-hsiRNA\(^{NTC}\)-injected animals is likely due to variability in tumor size at the two-week time point. Taken together, these data suggest that the Chol-hsiRNAs chemical architecture is an effective platform to silence gene expression in brain tumors \textit{in vivo} by direct intratumoral administration.

\textbf{Discussion}

GBM tumors are highly heterogeneous cellular ecosystems, composed of self-renewing stem-like cells (GSCs), differentiated cells, nonmalignant brain cells, and immune cells(23). GSCs are emerging as key effectors in tumor malignancy, resistance, and reoccurrence. Therapeutic targeting of GSCs represents one of the most direct routes towards curbing GBM tumorigenesis.
The outstanding challenges in targeting GSCs are identifying a therapeutic platform that is capable of modulating expression or activity of GBM driver genes and achieving targeted GSC delivery. Genome sequencing, transcriptomic and epigenetic profiling have unveiled gene clusters altered or misregulated in GBM, including aberrant methylation sites, histone modification patterns, and chromatin architectures(23,24). The vast majority of these targets are undruggable by conventional small molecule and biologics approaches(10). Oligonucleotides are powerful tools for precise, multi-target gene silencing in the central nervous system(11,15,25), as both potential therapeutics and as methods of probing the functional genomics of GBM(26). However, delivery is the key obstacle preventing immediate deployment of siRNAs against the molecular drivers of GBM.

The two primary approaches to achieve synthetic siRNA delivery are through formulations or direct conjugation(27). siRNA formulations or encapsulations (e.g. lipid nanoparticles) that are typically used to improve stability and retention in other organs show pronounced neuroinflammation and neuronal toxicity in brain(28-30). Direct lipid conjugation holds great promise to achieve safe, carrier-free siRNA delivery for neurological applications(11,15,25). Previously, we described the pharmacokinetic properties of cholesterol-conjugated hsiRNAs in mouse brain, demonstrating that Chol-hsiRNAs were rapidly internalized into neurons and glia, inducing potent gene silencing after an intraparenchymal injection in mouse striatum(11). Due to their lipophilicity, Chol-hsiRNAs were retained around the site of injection, exhibiting highly efficient cellular internalization(11). These considerations prompted us to investigate Chol-hsiRNAs as a platform for localized RNAi-mediated gene silencing in GSC-derived established tumors.
In the current report, we demonstrate that Chol-hsiRNAs are rapidly internalized into patient-derived GBM8 neurospheres, inducing significant mRNA and protein silencing in vitro with no observable toxicity. We show that Chol-hsiRNAs distribute broadly throughout established GBM8 brain tumors, inducing potent (>90%) and durable (7 d) silencing of tumor-specific reporter genes. This comprises the first report of efficient gene silencing in a GSC-derived orthotopic brain tumor by a lipid-conjugated siRNA. Chol-hsiRNAs have predictable pharmacokinetic properties and can be used to target single genes, or used in combination to modulate entire networks(9). Moreover, the low dosing requirements afforded by chemically modified siRNAs with enhanced stability and efficiency minimize the risk of off-target effects, collateral damage to healthy tissues, and unintended systemic toxicity(9). For instance, the most clinically advanced siRNA (a fully 2’ substituted, trivalent N-acetylgalactosamine-siRNA conjugate) achieved over six months of sustained target silencing in patients after a single dose, with no serious adverse events(31).

Recent key studies have validated multiplexed siRNA therapies as a strategy for improving survival in GSC tumor models(10), and have demonstrated the potential for unformulated oligonucleotides to effectively penetrate the blood-tumor barrier at high doses(20,32). We continue to investigate the ability of other classes of lipid-conjugated siRNAs to achieve potent gene silencing in brain tumors through systemic or CSF delivery.

Polyunsaturated fatty acid-siRNAs exemplify a new class of conjugates we are exploring for gene silencing in CNS (15,25). Last year witnessed the clinical approval of Spinraza (nusinersen), a splice-switching antisense oligonucleotide for the treatment of spinal muscular atrophy(33). The rapid evolution of oligonucleotide chemistries will soon yield entities capable
of translation into potent new GBM therapies, based on our expanding knowledge of the genetic changes that drive the biogenesis and growth of these devastating tumors.

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References


**Figure Legends**

**Figure 1.** Human primary GBM8 neurospheres productively internalize cholesterol-conjugated hsiRNAs (Chol-hsiRNA) in vitro. (a) Molecular model of a fully chemically modified, hydrophobic siRNA (Chol-hsiRNA) containing a cholesterol-tetraethyleneglycol (TEG) linker (blue) with 2’ fluoro (gray), 2’-O-methyl (black), phosphorothioate (red), and 5’-phosphate (purple) stabilizing modifications (PyMOL, Version 1.8, Schrödinger) (b) Internalization kinetics into GBM8 neurospheres were assessed using 0.75 μM Cy3-Chol-
hsiRNA<sub>His</sub> for 0–5 h. Nuclei (Hoechst), blue; Chol-hsiRNA (Cy3), red. Intensity and exposure settings were kept consistent between samples. Silencing of (c) <i>PPIB</i> and (d) <i>HTT</i> mRNA in GBM8 neurospheres following treatment with Chol-hsiRNA<sup>PPIB</sup> or Chol-hsiRNA<sup>HTT</sup>, respectively. <i>PPIB</i> and <i>HTT</i> mRNA levels were measured after 72 h and normalized to a housekeeping gene (<i>HPRT</i>), and represented as percentage of untreated control (n = 3 biological replicates, mean ± SD). UNT – untreated cells, NTC – non-targeting (scrambled) Chol-hsiRNA control.

**Figure 2. Firefly luciferase-targeting Chol-hsiRNA silences f<i>Luc</i> mRNA and protein expression in GBM8 neurospheres.** (a) Human primary GBM8 cultures were incubated with Chol-hsiRNA<sup>f<i>Luc</i></sup> at concentrations shown and <i>fLuc</i> mRNA levels were measured at 72 h and normalized to the human housekeeping gene <i>PPIB</i>. <i>fLuc</i> mRNA levels are represented as percentage of untreated control (n = 3, mean ± SD). (b,c) GBM8 cultures were incubated with (b) Chol-hsiRNA<sup>f<i>Luc</i></sup> or (c) Chol-hsiRNA<sup>NTC</sup> at concentrations shown for 96 h. Luciferase activity is represented as percent of untreated control (n = 3 biological replicates, mean ± SD). UNT – untreated cells, NTC – non-targeting siRNA control. (d) GBM8 neurospheres were monitored for number, size, and morphology following incubation with untreated medium, 3 µM Chol-hsiRNA<sup>f<i>Luc</i></sup>, and 6 µM Chol-hsiRNA<sup>NTC</sup>. Representative images are shown.

**Figure 3. Chol-hsiRNA distributes evenly throughout the brain tumor xenograft after intratumoral injection.** Cy3-labeled Chol-hsiRNA<sup>HTT</sup> was injected into established orthotopic GBM8 brain tumors. Distribution was analyzed at 24 h post-injection. (a) Tiled (10X) and magnified (20X) coronal images of H&E staining delineating tumor border in xenografted lobe.
(b) Tiled (10X) and magnified (40X) fluorescence images of coronal brain sections imaged for nuclear (DAPI, blue) and Chol-hsiRNA (Cy3, red) signal. White arrows indicate Chol-hsiRNA aggregation with axonal bundles. Yellow arrows indicate Chol-hsiRNA accumulation in cytoplasmic foci. Representative images, confirmed in two separate experiments.

**Figure 4. Chol-hsiRNA silences human HTT mRNA and firefly luciferase protein expression in brain tumors after direct injection.** (a) Changes in human HTT mRNA levels one week after intratumoral injection of aCSF, 5 nmol of non-targeting control Chol-hsiRNA$^{NTC}$ (NTC), 5 nmol Chol-hsiRNA$^{HTT}$, or ICV injection of 5 nmol Chol-hsiRNA$^{HTT}$ into four-week-old GBM8 tumors. Data was normalized for the human housekeeping gene PPIB, and represented as percent of untreated control ($n = 5$ mice, mean ± SD, three biopsies per striatum) (b) Changes in tumor-associated luciferase activity one week after intratumoral injection of Chol-hsiRNA$^{Fluc}$ (2.5 or 5 nmol) or Chol-hsiRNA$^{NTC}$ (5 nmol) in two-week-old GBM8 tumors ($n = 5$ mice, mean ± SD, three biopsies per striatum). Percentages reflect degree of luminescence reduction. ($^*P<0.05$, $^{**}P<0.01$, $^{***}P<0.001$, $^{****}P<0.0001$).
a) Implant GBM8 cells into right striatum (n = 26)
Monitor tumor growth by IVIS
Inject hsiRNA (mRNA)
Collect tissue

T = 0 days 14 d 28 d 34 d

Human H1T mRNA Expression, (% Control)

CSF NTC IT ICV

Chol-hsiRNA$^{HTT}$ (5 nmol)

45% *** 23% *

b) Implant GBM8 cells into right striatum (n = 26)
Inject hsiRNA (protein)
Collect tissue

T = 0 days 14 d 21 d

Firefly Luciferase Luminescence (RLU $10^5$)

CSF NTC 2.5 nmol 5 nmol

Chol-hsiRNA$^{flu}$ (intratumoral)

89% **** 90% ****

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