A nuclear function for an oncogenic microRNA as a modulator of snRNA and splicing

Rachid El Fatimy1,2†, Yanhong Zhang1†, Evgeny Deforzh3, Mahalakshmi Ramadas1, Harini Saravanan1, Zhiyun Wei1,3, Rosalia Rabinovsky1, Nadiya M. Teplyuk1, Erik J. Uhlmann1 and Anna M. Krichevsky1*

Abstract

Background: miRNAs are regulatory transcripts established as repressors of mRNA stability and translation that have been functionally implicated in carcinogenesis. miR-10b is one of the key onco-miRs associated with multiple forms of cancer. Malignant gliomas exhibit particularly striking dependence on miR-10b. However, despite the therapeutic potential of miR-10b targeting, this miRNA's poorly investigated and largely unconventional properties hamper the clinical translation.

Methods: We utilized Covalent Ligation of Endogenous Argonaute-bound RNAs and their high-throughput RNA sequencing to identify miR-10b interactome and a combination of biochemical and imaging approaches for target validation. They included Crosslinking and RNA immunoprecipitation with spliceosomal proteins, a combination of miRNA FISH with protein immunofluorescence in glioma cells and patient-derived tumors, native Northern blotting, and the transcriptome-wide analysis of alternative splicing.

Results: We demonstrate that miR-10b binds to U6 snRNA, a core component of the spliceosomal machinery. We provide evidence of the direct binding between miR-10b and U6, in situ imaging of miR-10b and U6 co-localization in glioma cells and tumors, and biochemical co-isolation of miR-10b with the components of the spliceosome. We further demonstrate that miR-10b modulates U6 N-6-adenosine methylation and pseudouridylation, U6 binding to splicing factors SART3 and PRPF8, and regulates U6 stability, conformation, and levels. These effects on U6 result in global splicing alterations, exemplified by the altered ratio of the isoforms of a small GTPase CDC42, reduced overall CDC42 levels, and downstream CDC42-mediated effects on cell viability.

Conclusions: We identified U6 snRNA, the key RNA component of the spliceosome, as the top miR-10b target in glioblastoma. We, therefore, present an unexpected intersection of the miRNA and splicing machineries and a new nuclear function for a major cancer-associated miRNA.

Keywords: miR-10b, Glioblastoma, Nucleus, U6 snRNA, Splicing machinery, CDC42

Background

miR-10b is one of the miRNAs most strongly associated with various types of cancers [1, 2]. Since the initial report of the aberrant miR-10b expression and association with breast cancer metastasis in 2007 [3], multiple studies have linked miR-10b to cancer progression, regulation of specific cancer-promoting signaling pathways, and clinical outcomes in a broad spectrum of malignancies [4]. miR-10b has been since implicated in cancer cell proliferation, epithelial-mesenchymal transition, invasion, metastasis, angiogenesis, and drug resistance. In addition, an investigation of miR-10b-deficient mice demonstrated that this miRNA is essential in tumorigenesis [5]. These data led to active exploration of

*Correspondence: akrichevsky@bwh.harvard.edu
†Rachid El Fatimy and Yanhong Zhang contributed equally to this work.
1 Department of Neurology, Brigham and Women’s Hospital and Harvard Medical School, 60 Fenwood Rd, Room 9002T, Boston, MA 02115, USA
Full list of author information is available at the end of the article
miR-10b as a therapeutic target as well as a cancer biomarker by academia and the biotechnology industry.

Perhaps the most distinct and intriguing is the activity of miR-10b in malignant gliomas, primary brain tumors affecting both adult and pediatric populations. While in most other cancers miR-10b primarily drives metastasis, in gliomas it emerged as a critical survival factor required for tumor cell viability [6]. The major difference in miR-10b activity in cancer cells of different lineages likely stems from its expression patterns: whereas expressed in normal extracranial tissues, miR-10b gene is completely silenced in the fetal and adult brain cortex but deregulated in the low- and high-grade gliomas [6, 7]. This aberrant out-of-place expression and activity of miR-10b in the brain tumors become addictive for the tumor cells due to the aberrant out-of-place expression and activity of miR-10b [8].

miR-10b inhibiting (CLEAR-CLIP) [14], led to the discovery of U6 small nuclear RNA (snRNA), the core component of a spliceosome as the major miR-10b binding transcript. We further developed a range of complementary biochemical and imaging-based techniques to investigate a new and unanticipated miRNA function in controlling U6 snRNA metabolism, spliceosome assembly, and alternative splicing in the cancer cells. Therefore, this study further expands the network of miRNA regulation beyond miRNA and IncRNA targets and demonstrates the interface of miRNA and splicing machineries.

Methods

Cell culture
LN229, U251 (human glioblastoma), MDA-MB-231 (human breast cancer), HEK-293 (human embryonic kidney), and Hela cell lines were cultured in Dulbecco’s Modified Eagle Media (DMEM) supplemented with 10% Fetal Bovine Serum (FBS) (WISENT), 1% penicillin and streptomycin (Gibco) and passaged by trypsinization (Corning®). SH-Sy-5y (human neuroblastoma) cell line was purchased from ATCC and passaged in DMEM-F12 supplemented with 10% FBS, 1% MEM Non-Essential Amino Acids Solution (NEAA) (Gibco), 1% penicillin and streptomycin (Gibco). Low-passage GSC (GBM4 and GBM8) cells were provided by Dr. Hiroaki Wakimoto (MGH) and cultured as neurospheres in Neurobasal-medium (Gibco) supplemented with 0.5% N-2 (Gibco), 2% B-27 (Gibco), 1.5% glucose (Gibco), 0.02 ng/ml epidermal growth factor (EGF) (Gibco), and 20 ng/ml fibroblast growth factor (FGF) (PeproTech) in ultralow attachment flasks or plates. GSC neurospheres were passaged using the NeuroCult chemical dissociation kit (Stemcell Technologies). Human astrocytes were cultured in astrocyte medium (ScienCell Research Laboratories) supplemented with 2% FBS, 1% antibiotic solution and 1% astrocyte growth supplement (ScienCell Research Laboratories) and passaged using cell/tissue dissociation solution (Innovation Cell Technologies, Inc.). All cells were maintained at 37°C and 5% CO₂.

Oligonucleotides, siRNAs, and cell transfection

A chemically modify 2′-O-methoxymethyl ASO inhibitor of miR-10b (5′-CACAATCGTTTCAGGGTA-3′, miR-10b ASO) and control oligonucleotide of the same chemistry (5′-ACATACTCCTTTCTACAGTGCA-3′) were designed as previously reported [7]. The following U6 ASOs and corresponding control have been designed based on the [15, 16]: U6 ASO #1 mUmGmGmAmACG CTTACAGAATtmUmGmCmG, U6 ASO #2 mUmGmAmAmAAGACACAGATCTACAmCmUmGmA; control mCmCmAmGmGCTACCACAmCmUmU (m=2′-O-methyl, with phosphorothioate backbone). MiR-10b mimic (cat no: C-300550-07-0005) and miR-10b as a therapeutic target as well as a cancer biomarker by academia and the biotechnology industry.

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matching inhibitors or mimics (final concentration of 50 nM), the next day, the cells were transfected with either miR-U6 ASOs (final concentration of 1 nM), CDC42 siRNAs (GTG GAG TCC GTA-3; CDC42-isoform2-specific siRNA (AGC CUC CAG CA-3), CDC42-isoform1-specific siRNA (AGC CUC CAG CA-3), CDC42-isoform2-specific siRNA (AGC CUC CAG CA-3), and control siRNA (UGG UUU ACA UGU CUA UGU A), purchased from Santa Cruz Biotechnology (sc-29,256). The CDC42 overexpression and control plasmids were obtained from OriGene. For transfections, adherent LN229 and U251 cells were plated in 24 well plates (Corning®) at 2 × 10^4/well. On the next day, the cells were transfected with either miR-10b inhibitors or mimics (final concentration of 50 nM), U6 ASOs (final concentration of 1 nM), CDC42 siRNAs (final concentration of 100 nM), or corresponding control oligonucleotides using lipofectamine 2000 (ThermoFisher Scientific). For GSC transfection, neurospheres were dissociated to single cell suspension (Stemcell Technologies) and transfected with the oligonucleotides using Neuromag 500 (OZ Biosciences).

RNA isolation and quantitative RT-PCR (qRT-PCR)
RNA was extracted using the Total RNA Purification Kit (Norgen Biotek, Catalogue # 17250) and quality of the samples examined using Nanodrop. For microRNA analysis, 200 ng of the total RNA was reverse transcribed using miRCURY LNA RT kit (QIAGEN, 339340). The cDNA libraries were then diluted 80 times and used for the analysis of gene expression by qPCR using miRCURY LNA™ primers (YP00203907 for U6; YP00205637 for miR-10b, YP00204339 for miR-125a) (QIAGEN). The cells were treated with Actinomycin D (final concentration 5 μg/ml, in 0.05% DMSO) (Sigma) for 2-36 h, followed by the RNA isolation and analysis. To examine the effects of miR-10b on U6 stability, cells were transfected with either miR-10b inhibitor or mimic and matching control oligonucleotides and culture media was replaced with the media containing 5 μg/ml of Actinomycin D at 24 h after transfections. RNA was isolated at different time points and qRT-PCR reactions were performed as described above.

RNA stability and half-life
For monitoring half-life of miR-10b and U6 snRNA, glioma LN229 and U251 cells were cultured in 6-well plates. The cells were treated with Actinomycin D (final concentration 5 μg/ml, in 0.05% DMSO) (Sigma) for 2-36 h, followed by the RNA isolation and analysis. To examine the effects of miR-10b on U6 stability, cells were transfected with either miR-10b inhibitor or mimic and matching control oligonucleotides and culture media was replaced with the media containing 5 μg/ml of Actinomycin D at 24 h after transfections. RNA was isolated at different time points and qRT-PCR reactions were performed as described above.

Western blot
Cells were harvested using RIPA buffer (Boston Bio-Products) and proteins were quantified using Micro BCA™ Protein assay kit (ThermoFisher Scientific). The cell lysates containing equal amounts of total proteins (30 μg) were separated by SDS PAGE in 4-12% Bolt Bis-Tris Plus gels (ThermoFisher Scientific) and transferred to 0.2 μm Immobilon™ PVDF membrane (BIO-RAD). After blocking with 5% (w/v) fat free milk in phosphate buffer saline (PBS) with 0.1% Tween-20 (PBS-T), membranes were incubated with primary antibodies (anti-HSP90 (Cell Signaling Technology, #4879S, 1:3000 dilution); anti-Lamin B (ABclonal, Kit, NB1001). Briefly, total RNA (300 ng) was denatured at 70°C for 5 min and resolved on 4-20% Tris-borate-EDTA (TBE) Urea gel (ThermoFisher scientific) in cold 0.5 × TBE buffer (Invitrogen) at 60 V for 5-6 h. The RNA was then transferred to a nylon membrane at 60 V in 0.5 × TBE with 20% methanol for 1 h at 4°C. The RNA was crosslinked to the membrane by UV irradiation (120 mJ/cm²). The membranes were incubated with biotin-prelabeled probes overnight at 42°C and rinsed with hybridization washing buffer for 30 min at 42°C. After blocking, the membranes were incubated with Streptavidin-HRP conjugate and signals developed using a chemiluminescence imaging system. The oligonucleotide probes for Northern blots were as following: U2 (ACACCTTATGTTAGC AAAAGGGCCGAGAAGCGAT); U4 (GCC AATGCCGACTATATTG); U6 (AGTATATGTGCT GCCGAAGCGAGCACT). For the analysis of native RNA conformation, cells were washed and resuspended in a mild lysis buffer (1% Igepal, 0.5% sodium deoxycholate, 0.1% SDS), and deproteinized in a buffer containing 1 mg/ml proteinase K, 50 mM NaCl, 10 mM EDTA, and 0.5% SDS for 30 min at 37°C. Total RNA (5 μg) was separated in 4-20% TBE gel (ThermoFisher scientific), transferred to a nylon membrane, cross-linked, and the hybridizations were performed as described above.

Denaturing and native northern blot
RNA samples were analyzed by denaturing Northern blotting, following the manufacturer’s instructions (Signosis, High Sensitive MiRNA Northern Blot Assay Kit, NB1001). Briefly, total RNA (300 ng) was denatured at 70°C for 5 min and resolved on 4-20% Tris-borate-EDTA (TBE) Urea gel (ThermoFisher scientific) in cold 0.5 × TBE buffer (Invitrogen) at 60 V for 5-6 h. The RNA was then transferred to a nylon membrane at 60 V in 0.5 × TBE with 20% methanol for 1 h at 4°C. The RNA was crosslinked to the membrane by UV irradiation (120 mJ/cm²). The membranes were incubated with biotin-prelabeled probes overnight at 42°C and rinsed with hybridization washing buffer for 30 min at 42°C. After blocking, the membranes were incubated with Streptavidin-HRP conjugate and signals developed using a chemiluminescence imaging system. The oligonucleotide probes for Northern blots were as following: U2 (ACACCTTATGTTAGC AAAAGGGCCGAGAAGCGAT); U4 (GCC AATGCCGACTATATTG); U6 (AGTATATGTGCT GCCGAAGCGAGCACT). For the analysis of native RNA conformation, cells were washed and resuspended in a mild lysis buffer (1% Igepal, 0.5% sodium deoxycholate, 0.1% SDS), and deproteinized in a buffer containing 1 mg/ml proteinase K, 50 mM NaCl, 10 mM EDTA, and 0.5% SDS for 30 min at 37°C. Total RNA (5 μg) was separated in 4-20% TBE gel (ThermoFisher scientific), transferred to a nylon membrane, cross-linked, and the hybridizations were performed as described above.

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A16685, 1:1000 dilution); anti-CDC42-iso1 (Millipore, ABN1646, 1:1000 dilution); anti-CDC42-iso2 (07-1466, Millipore, 07-1466, 1:1000 dilution); anti-CDC42-total (PA1-092, ThermoFischer Scientific, PA1-092, 1:1000 dilution); α-Tubulin (Sigma, T9026, 1:500 dilution) at 4°C overnight. The membranes were further washed with PBS-T buffer and incubated with secondary antibodies (Cell Signaling Technology, #14708 or #14709, 1:10,000 dilution) for 2h at room temperature. The blots were developed using ECL reagent Supersignal West Pico chemiluminescent Substrate (ThermoFisher Scientific).

Cell fractionation and isolation of nuclear fractions

For subcellular fraction, 5 × 10⁶ cells were collected and washed twice with PBS and transferred to a microcentrifuge tube. The pellets were re-suspended in 1x hypotonic buffer (20 mM Tris-HCl (pH = 7.4), 10 mM NaCl, 3 mM NaCl₂, 1 mM Triton X-100, 1 mM EDTA, 10% glycerol, 1 mM EGTA, 0.1% SDS, 1 mM NaF, 0.5% deoxycholate, 20 mM Na₃P₂O₇) supplemented with 1 mM PMSF and Protease Inhibitor Cocktail (Sigma), vortexed, and centrifuged again. The resulting supernatants contained soluble nuclear fractions and the pellets containing nuclear fractions have been collected. The pellets were resuspended in the cell extraction buffer (10 mM Tris (pH = 7.4), 2 mM Na₃VO₄, 100 mM NaCl, 1% Triton X-100, 1 mM EDTA, 10% glycerol, 1 mM EGTA, 0.1% SDS, 1 mM NaF, 0.5% deoxycholate, 20 mM Na₃P₂O₇) supplemented with 1 mM PMSF and Protease Inhibitor Cocktail (Sigma), vortexed, and centrifuged again. The resulting supernatants contained soluble nuclear fractions and the pellets contained nuclear insoluble fractions. The isolated nuclear fractions were subjected to ultra-centrifugation on the linear glycerol gradient (from 5 to 60%) for 2 h at 4°C, 26,000 rpm using SW55 rotor (Beckman Coulter L-90 K optima), and aliquoted to 200 μl fractions [17]. These fractions have been used for RNA and protein isolation and analysis.

Immunofluorescence

Cells were washed with PBS and fixed with 4% paraformaldehyde (PFA), permeabilized with 0.5% Triton X-100 in PBS, and blocked in 5% BSA for 1h. The cells were then incubated with primary antibodies (rabbit anti-KI-67, Abcam, ab16667, 1:250 dilution) overnight at 4°C and further incubated with secondary antibodies (goat anti-rabbit Alexa fluoro 488, Invitrogen, 1:200 dilution). The nuclei were stained with DAPI (1:1000). The staining was visualized and imaged using a scanning confocal microscopy.

Fluorescence in-situ hybridization (FISH) and immunofluorescence

RNA FISH was performed using a ViewRNA cell Plus Assay Kit (Invitrogen, #88-1900-99) following the manufacturer instructions. Briefly, 1 × 10⁶ cells per well were cultured in a chamber slide (8 Chamber Polystyrene Vessel Tissue Culture Treated Glass Slide, FALCON, #354118) and fixed with the fixation solution for 1h at room temperature. For the tumor analysis, resected GBM specimens were obtained from the Brigham and Women’s Hospital, per a protocol approved by the Institutional Review Board. Frozen human kidney tissue was purchased from BioChain (Cat no: # T1234142). Frozen GBM tissues were fixed with 4% PFA overnight at 4°C and blocked in 30% sucrose for 4-6h. After transferring to the optimal cutting temperature compound (OCT), GBM tissues were cut into 15 μm-thick sections and placed into acetone for 5 min at −20°C. After three washes with RNase inhibitor-containing PBS, the cells/tissue slides were permeabilized with 0.3% Triton-X 100 in PBS for 5 min on ice. The cells/tissue slides were hybridized with probes specific for mature miR-10b, miR-21, or U6 RNA (Invitrogen, 1:100 dilution) or pri-miR-10b (Biosearch Technologies, 1:100 dilution) for 2h at 40°C. To control for background staining, the probes were omitted in parallel experiments. The cells were further washed with wash buffer; the tissue slides were washed with 2X SSC for 5 times at 40°C. The cells/slides were hybridized sequentially with PreAmplifier, Amplifier and Label probe solution for 1h at 40°C, stained with 1 × DAPI (4′,6-diamidino-2-phenylindole), mounted in a mounting media (ThermoFisher Scientific), and observed under a confocal microscope (CZ Scan). For the FISH combined with Immunofluorescence, cells were fixed in 4% PFA, treated with 0.3% Triton-X 100 solution, and blocked with 5% BSA. Tissues slides were placed into acetone and then treated with antigen repair solution (20 mg/ml Proteinase K, 1 M Tris-HCl (pH = 8.0), 0.5 M EDTA (pH = 8.0)) for 5 min at 37°C. The step of overnight incubation at 4°C with primary antibody (anti-PRPF8, ABclonal, A6053, 1:200 dilution; or anti-SART3, GeneTex, GTX107684, 1:200 dilution) was included before hybridization with FISH probes and the subsequent signal amplification steps. The probes were then washed and the cells/slides incubated with secondary antibodies (goat anti-rabbit Alexa fluoro 488, Invitrogen, 1:200 dilution) for 1h at room temperature. They were further washed with PBS three times, stained with 1 × DAPI, and images taken using confocal microscopy. Quantification of the colocalization was determined by Image J pro 6.0.
CLEAR-CLIP: covalent ligation of endogenous Argonaute bound RNA - crosslinking and immunoprecipitation

CLEAR-CLIP was performed according to the published protocol with minor modifications [14]. Briefly, LN229 cells were irradiated with UV-C light (2 times by 2400 J/m²) to covalently cross-link proteins to nucleic acids. The cells were lysed, treated with DNase (Promega), followed by the partial RNA fragmentation using low concentrations of the RNase I (0.002 U/ml, 5 min), and treatment with the RNase inhibitor (RNAsin Plus at 0.5 U/µl) to quench RNase activity. The miRNA–RNA complexes were immunopurified using the anti-Ago-2 antibody (Millipore, #03-110) immobilized on immunoglobulin G-coated magnetic beads (Life Technologies). The beads were washed, treated with T4 Polynucleotide Kinase (PNK, with no 3’-phosphatase activity) (NEB, M0236S), and miRNA–RNA chimeras were produced with T4 RNA Ligase 1 (NEB, M0204S). The beads were further washed and subjected to alkaline phosphatase treatment (Roche, #10713023001) to remove 3’ phosphate. The 3’ linker (5’-rAppGTGTCAAGTTCCAGCG-3’, Dharmacon) was added using T4 RNA Ligase 2 (NEB, M0242S), followed by the PNK (NEB, M0201S) treatment. The samples were resolved on 8% acrylamide gel (ThermoFisher Scientific), transferred to nitrocellulose membrane and RNA eluted from the membrane as described [18]. Briefly, the membranes were cut into small pieces of 0.5–2 mm, transferred to 1.5 mL low-binding tubes, and incubated in the solution containing 100 mM Tris pH 7.4, 50 mM NaCl; 10 mM EDTA, and 4 mg/ml Proteinase K for 20 min at 55 °C with constant mixing at 1200 rpm.

To covalently crosslink the proteins to nucleic acids, glioma cells (2 × 10⁷) were subjected to UV irradiation (200 ml/cm²). The cells were then lysed with RIPa lysis buffer containing Protease Inhibition Cocktail and RNase Inhibitor followed by DNase treatment (Promega). The immunoglobulin-coated protein magnetic beads (ThermoFisher scientific) were incubated with antibodies (anti-pseudouridine, MBL, D347-3; anti-N6-methyladenosine (m6A), Millipore, MABE1006; anti-PRPF8, Abclonal, A6053; anti-SART3, GeneTex, GTX107684; anti-Ago-pan, Millipore, MABE56; anti-Ago2, Millipore, 03-110; anti-LSM8, Invitrogen, PA5-69098; anti-LSM4, Abclonal, A5891; anti-DHX8, Abclonal, A14724; anti-RBM22, Abclonal, A10025; anti-SNRNP200, Abclonal, A6063; anti-LMN1, Abclonal, A16685) to immunopurify the respective RBP complexes. The complexes were washed with RIP buffer, and the samples treated with proteinase K buffer (10% SDS and 10 mg/ml proteinase K in RIPA buffer) for 30 min at 37 °C with shaking. RNA was further isolated using phenol:chloroform:isoamyl alcohol (25:24:1) solution, precipitated using isopropanol, and suspended in RNase-free water.
Library constructions, RNA sequencing, and bioinformatics analysis

GBM4 cells were transfected with miR-10b inhibitor, U6 ASO #1, and corresponding control oligonucleotides for 72h. The libraries have been constructed and sequenced by Novogene Corporation. Briefly, a total amount of 500 ng RNA per sample was used as input material. Ribosomal RNA was removed by Epicentre Ribo-zero™ rRNA Removal Kit (Epicentre, USA), and the libraries have been generated using NEBNext® Ultra™ Directional RNA Library Prep Kit for Illumina® (NEB, USA). The libraries have been purified using the AMPure XP system, and their quality assessed by Agilent Bioanalyzer 2100. The libraries have been sequenced on an Illumina platform and at least 12G of paired-end reads per sample were generated (Novogene). The data filtering included removing adaptors, removing reads containing \(N\) > 10% (\(N\) represents base that could not be determined), the Qscore (Quality value) of over 50% bases of the read is \(<=\) 5. Alternative splicing analysis was performed by the software rMATS, a statistical method for robust and flexible detection of differential alternative splicing from replicate RNA-Seq data. It identifies alternative splicing events corresponding to all major types of alternative splicing patterns and calculates the \(P\) value and FDR for differential splicing. These types include exon skipping (SE), alternative 5′ splice sites (A5SS), alternative 3′ splice sites (A3SS), mutually exclusive exons (MXE), and retained introns (RI).

Luciferase reporter assay

Full-length human RNU6 sequence (110 nucleotides) was amplified by PCR and cloned into psiCHECK2 (Promega), downstream of Renilla luciferase, using XhoI and NotI restriction sites. Glioma cells were transfected with 100 ng of the reporter constructs, including the original “empty” plasmid and U6 reporter, and 24h later transfected with either miR-10b mimic (final concentration of 25 nM), miR-10b inhibitor (final concentration of 50 nM), or the corresponding control oligonucleotides. Luciferase luminescence was measured using the Dual-Glo luciferase assay system (Promega, E2920) 24h after the last set of transfections.

Statistical analysis

Details pertaining to all statistical analyses can be found in the figure legends.

Results

miR-10b binds to snRNA U6 in glioma cells

To investigate endogenous miR-10b targets, we employed the modified CLEAR-CLIP protocol in glioma cells [14]. The technique relies on miRNA crosslinking with proximal transcripts, ligation, and sequencing of chimeras. Therefore, it provides an unbiased mapping of the miRNA-target interactions, independent of bioinformatic predictions (Fig. 1a). The identified miR-10b chimeras contained fragments matching multiple predicted and validated mRNA targets (Fig. 1b and Supplementary Table 1). Of the identified miR-10b-binding transcripts, 115 corresponded to protein-coding and 31 to non-protein coding genes, and the majority did not contain miR-10b seed, despite the extensive base-pairing with miR-10b (Fig. 1c and Supplementary Fig. 1 and Supplementary Table 2). Surprisingly, the most common chimeras were formed by miR-10b and the sequences corresponding to U6 snRNA. Inspection of U6 snRNA transcript demonstrated extensive base pairing between its 3′ end and miR-10b, with a stretch of 10 complementary nucleotides at the positions 14-23 of miR-10b, and additional complementarity beyond this site (Fig. 1d). To validate the binding between miR-10b and U6, we further used CLEAR-CLIP to produce miRNA chimeras from several miR-10b-expressing cancer cell lines (Supplementary Fig. 2) and detect the product with a pair of PCR primers, one corresponding to miR-10b and another to U6. We detected Ago-bound miR-10b- U6 chimeras in glioma cell lines and GSCs, with trace levels also present in other cancerous non-glioma cells, but not in IgG control groups (Fig. 1e). Collectively, these data demonstrate that miR-10b binds to U6 snRNA in various cancer cells, and this intermolecular interaction is prevalent in glioma cells.

snRNA U6 is a major RNA component of the spliceosome. It localizes predominantly, if not exclusively, to the nuclear space [22]. Mature miRNAs are thought to operate mostly in the cytosolic compartment; however, in specific conditions some of them could be found in the nucleus [23, 24]. In agreement with this, in glioma cells, the major enzymatic complex responsible for the final step of miRNA processing, Dicer, is aberrantly localized to the nucleus [12]. To investigate miR-10b subcellular localization, we fractionated glioma cells into cytosolic and nuclear fractions. More than half of the cellular miR-10b pool was confined to the nuclei; of this nuclear pool, >40% was associated with the insoluble fractions containing the spliceosomal complexes (Fig. 2a).

To further explore the association between miR-10b and snRNA U6 in cancer cells, we visualized mature miR-10b and U6 in LN229 and U251 glioma cells and other cancer cells using RNA in situ hybridization (FISH). miR-10b localized in multiple nuclear and perinuclear speckles and often co-localized with U6 snRNA (Fig. 2b). The co-localization was more common in glioma than in other cell lines (Fig. 2b). Importantly, patient-derived GBM tumors also exhibited remarkable, in some cells...
almost perfect colocalization between miR-10b and U6 (Fig. 2c). No colocalization was detected between U6 and another highly abundant cancer-associated miRNA, miR-21 (Fig. 2c), as well as with miR-10b close paralogue miR-10a and the pri-miR-10b transcript (Supplementary Fig. 3a, b), indicating the selectivity and sequence-specificity of the U6 binding to miR-10b in glioma. Of note, normal kidney tissues, expressing relatively high miR-10b...
Fig. 2 (See legend on previous page.)
levels [25], exhibited minimal, if any, colocalization between the miR-10b and U6 (Supplementary Fig. 3c).

miR-10b is a part of the spliceosomal SART3 and PRPF8 RNPs

To further validate the binding between miR-10b and U6, we complemented the pool-down of Ago complexes with the similar RNA iCLIP technique employing various U6-interacting spliceosomal proteins as tags. Both U6 and miR-10b have been significantly enriched in the spliceosomal SART3 and PRPF8 complexes (Fig. 3a and Supplementary Fig. 4a, b). Furthermore, miR-10b- U6 chimeras have been detected in the cross-linked and ligated SART3 and PRPF8 complexes, similar to the Ago complexes (Fig. 3b) indicative of the miR-10b interaction with U6 snRNP in the splicing complex. SART3 or the spliceosome-associated factor 3 (aka hPrp24) is a core spliceosomal factor and a single spliceosomal protein that binds U6 selectively [26]. It participates in the U6 biogenesis, function in splicing (unwinding the U6 internal stem-loop, ISL, and annealing of U4 to U6), and recycling [27]. Prp8 or Pre-mRNA-processing-splicing factor 8, one of the largest and most conserved proteins in the spliceosome, is directly implicated in splicing fidelity [28]. Prp8 serves as a scaffold protein stabilizing the core of the spliceosome, which conformational rearrangement facilitates the 3D folding of the catalytically active U2/U6 RNA [29].

Using a combination of miRNA FISH and immunofluorescence (IF) techniques, we further visualized miR-10b in the nuclear SART3 speckles observed in glioma cells (Fig. 3c). miR-10b also partly colocalized with PRPF8 that was distributed in the nuclei more diffusely (Fig. 3d). Moreover, miR-10b colocalization with nuclear SART3 and PRPF8 RNPs was observed in patients-derived GBM tumors (Fig. 3e, f). Of note, miR-10b binds to the nucleotides 79-89 within the asymmetric bulge of U6, which is the SART3 binding site, suggesting the competitive manner of miR-10b and SART3 binding to U6. To test this hypothesis, we transfected glioma cells with the miR-10b antisense oligonucleotide (ASO) inhibitor and examined the effects of miR-10b KD on the U6 association with SART3 and PRPF8 using iCLIP. U6 association with both SART3 and PRPF8 was enhanced by the miR-10b inhibitor (Fig. 3g, h). Conversely, it was reduced by the synthetic miR-10b mimic (Fig. 3i, j). These results indicate that miR-10b interferes with U6 incorporation into SART3 and PRPF8 snRNPs. Of note, nuclear Ago2 was observed in glioma cells (Supplementary Fig. 5a) and both SART3 and PRPF8 were pulled-down by the Ago2 immunoprecipitation. Conversely, Ago2 and pan-Ago were immunoprecipitated in the SART3 and PRPF8 complexes (Supplementary Fig. 5b-e). These data further support the interference of the miRNA and the splicing machineries.

miR-10b regulates U6 snRNA levels, stability, modifications, and conformation

miRNAs are established regulators of gene expression that destabilize mRNA targets and reduce their translation. Here we investigated whether miR-10b regulates the levels of U6 snRNA. Two alternative techniques, qRT-PCR and Northern blot, demonstrated that miR-10b overexpression reduced U6 levels similarly to the effects induced by two alternative, fully complementary U6 ASOs (Fig. 4a-c). Conversely, miR-10b inhibitor upregulated the U6 levels (Fig. 4b, c). These effects were specific for U6 as other small RNAs were not affected by miR-10b modulators (Fig. 4c and Supplementary Fig. 6a). Furthermore, the experiments with transcription inhibitor actinomycin D demonstrated that miR-10b reduced the stability of U6, with miR-10b inhibitor and mimic causing stabilization and de-stabilization of U6, respectively (Fig. 4d and Supplementary Fig. 6c). These effects appeared independent of the conventional RISC machinery since a luciferase reporter with the inserted full-length U6 sequence has been affected by neither miR-10b inhibitor nor its mimic (Supplementary Fig. 6b).

To better understand how miR-10b regulates U6 stability, we investigated its effects on the selected U6 RNA modifications. U6 is pseudouridylated at several positions [30] and miR-10b binding to the U6 nucleotides 79-88 could involve the constitutively pseudouridylated U at position 86 (U86). It may also interfere with U31 pseudouridylation of the telestem and other U6 snRNA modifications and intramolecular secondary structures (Fig. 4a). Using iCLIP technique, we tested the effects of miR-10b modulation on the U6 pseudouridylation. miR-10b mimic strongly induced its levels whereas miR-10b inhibitor reduced it (Fig. 4e). Similar effects on the N-6-methyladenosine modification (m6A) of U6 found at position A43 have also been observed (Fig. 4e).

During the splicing cycle, U6 snRNA undergoes series of conformational changes required for accurate interactions with other snRNA and protein components of the spliceosome, recognition of mRNA splice sites and intron removal [30, 32]. We hypothesized that miR-10b binding affects U6 conformation and utilized native non-denaturing PAGE and Northern blot hybridization to test this hypothesis. Notably, transfection of glioma cells with both miR-10b mimic or inhibitor led to the appearance of a slower-migrating U6 variant, similar to one of the variants formed in the cells transfected with U6 ASO1 that also binds to the 3’ end of the U6, as does miR-10b (Fig. 4f). Of note, the patterns observed by non-denaturing Northern Blotting are indicative of conformation...
changes; however, the method lacks the resolution to claim that miR-10b mimic and inhibitor cause identical conformational alterations. The miR-10b mimic, whose sequence and chemistry copy the endogenous miR-10b, interferes with the 3′ end of U6 directly. The miR-10b inhibitor, however, more likely interferes with U6 via its
initial basepairing with miR-10b, probably forming a stab-
iling triple helix structure, and thereby extending U6 half-
life. Altogether, these data indicate that miR-10b
binding to U6 snRNA modulates U6 modifications, con-
formation, its binding to SART3 and PRPF8 protein part-
ers in the snRNP assembly, and overall U6 stability.

miR-10b is much shorter than U6 and is thought to be
less abundant than U6, raising the question of how it can
regulate U6. In contrast to snRNAs, miRNAs are among
the most stable cellular transcripts [33]. We, therefore,
postulated that a single copy of miR-10b could exert
its function on multiple U6 complexes. To directly com-
pare half-lives of the endogenous miR-10b and U6 in gli-
oma, we carried out the experiments using actinomycin
D. Notably, the U6 half-life is about 3-4h; however, miR-
10b exhibits much higher stability, with the half-life of at
least 36h (Fig. 4g and Supplementary Fig. 6d, e). These
results suggest that a single miR-10b molecule can bind
to and perturb the conformation and function of multiple
U6 snRNAs during its life cycle. To better understand the
stoichiometry of miR-10b and U6 binding in glioma, we
reanalyzed two large GBM RNAseq datasets for miR-10b
and U6 expression. This analysis demonstrated highly
variable levels of both miR-10b and U6 in GBM. Remark-
ably, their median levels were comparable in both data-
sets (Fig. 4h, i). Since miR-10b and U6 transcripts have
different length and modifications, the RNAseq data may
not accurately reflect their ratios; nevertheless, it strongly
suggests that miR-10b expression in glioma has been pre-
viously underestimated.

Interestingly, U6 levels in normal neuroglial cells,
including astrocytes and neuroprogenitors that are the
cells of glioma origin, are much lower than in glioma cells
(Supplementary Fig. 7a). This data is in agreement with
reports of 1) low U6 levels in the adult brain [15] and 2)
significantly higher U6 levels in certain carcinomas in
comparison to those in corresponding normal tissues
[34], and is plausible in view of the enhanced proliferation
and high transcriptional and splicing capacity of cancer
cells. Correspondingly, miR-10b overexpression in nor-
mal astrocytes had very strong effect on U6 levels (Sup-
plementary Fig. 7b). Considering that miR-10b is silenced
in normal neuroglial cells and strongly derepressed in
gliomas [6, 9], and miR-10b/ U6 ratio increases dramatic-
ally in glioma (Supplementary Fig. 7c), we propose that
at progressive stages of gliomagenesis and in diverse cell
subpopulations of highly heterogeneous astrocytomas,
the stoichiometry of miR-10b and U6 may change and
reach dynamic equilibrium.

miR-10b and U6 regulate alternative splicing of CDC42
miR-10b has been previously implicated in regulating
alternative splicing via targeting splicing factors such as
SRSF1 and MBNL1-3 [7, 35]. The observed direct
regulation of U6, the core component of the spliceo-
some machinery, however, suggests that miR-10b may
also have U6-mediated effects on splicing. The absolute
and relative levels of U6 in the spliceosome are highly
variable during development, across tissues, and cancer
samples, and U6 perturbations have profound impacts
on gene-specific differences in alternative splicing but
not transcriptome-wide splicing failure [15]. Consis-
tent with this, the RNA sequencing demonstrated global
changes in splicing patterns upon treatments of
GSCs with miR-10b inhibitor and U6 ASO (Fig. 5a and

(See figure on next page.)

**Fig. 4** miR-10b regulates U6 snRNA levels, stability, and conformation. a Sequence, putative secondary structure, nucleotide modifications of human U6, and miR-10b binding to U6 (top panel). The structure and modifications of human U6 have not been experimentally determined, and are shown to mimic that of yeast U6 [30]. Positions of U6-targeting ASOs are shown in the lower panel. U6 ASO-1 binds to the 3'end of U6, similarly to miR-10b. b qRT-PCR analysis of U6 levels in glioma cells and GSCs transfected with either U6 ASO, miR-10b inhibitor, mimic, or corresponding control oligonucleotides (mean ± SD, n = 3). P values were calculated using two-way ANOVA. e iCLIP with antibodies recognizing pseudouridylation (top) and m6A methylation (bottom) on LN229 and GBM8 cells transfected with either miR-10b inhibitor or mimic, and the corresponding control oligonucleotides, followed by U6 qRT-PCR detection, demonstrate that U6 modification are regulated by miR-10b (mean ± SD, n = 3). P values were calculated using two-tail unpaired t-test. f Representative native Northern blotting with U6-specific probe, demonstrates that U6 levels are regulated by miR-10b. The lower panel demonstrates that other small RNA species, resolved by denaturing electrophoresis, are not regulated by miR-10b. d qRT-PCR analysis of U6 levels in LN229 cells transfected with miR-10b inhibitor (top) or mimic (bottom), and treated with 5 μg/ml Actinomycin D demonstrates that miR-10b reduces U6 half-life (mean ± SD, n = 3). P values were calculated using two-way ANOVA. e iCLIP with antibodies recognizing pseudouridylation (top) and m6A methylation (bottom) on LN229 and GBM8 cells transfected with either miR-10b inhibitor or mimic, and the corresponding control oligonucleotides, followed by U6 qRT-PCR detection, demonstrate that U6 modification are regulated by miR-10b (mean ± SD, n = 3). P values were calculated using two-tail unpaired t-test. f Representative native Northern blotting with U6-specific probe demonstrates that miR-10b modulation leads to the appearance of an additional U6 structural variant. g Gioma cells treated with Actinomycin D, followed by qRT-PCR for miR-10b and U6 levels, exhibit high stability of miR-10b (mean ± SD, n = 3). P values were calculated using two-way ANOVA. h Expression of miR-10b and U6 in 155 GBM samples representing TCGA/GDC Pan-Cancer dataset, retrieved from UCSC XENA browser (https://xena.ucsc.edu/). The RNAseq data are FPKM normalized and presented as log2(FPKM-uq + 1). * P < 0.05; ** P < 0.01; *** P < 0.001; **** P < 0.0001
Supplementary Table 3, 4). Notably, alternative splicing of numerous genes changed similarly by both miR-10b inhibitor and U6 ASO1, with similarities observed in exon skipping (SE), alternative 5′ splice sites (A5SS), alternative 3′ splice sites (A3SS), mutually exclusive exons (MXE), and retained introns (RI) (Fig. 5a and Supplementary Table 3, 4).

To compare the effects of miR-10b and U6 perturbations, we utilized CDC42, one of the critical genes implicated in glioma biology, whose splicing is modulated by miR-10b [7]. There are two major CDC42 mRNA variants produced in GBM: the isoform ENST00000315554 (iso-1) and the predominant variant ENST00000344548 (iso-2) with the additional exon 2 and alternative last exon utilized (Fig. 5b). Although the ORFs of the two isoforms are very similar and translate to the proteins with only slightly altered C-terminus, their 3′ and 5′ UTRs are highly distinct. Iso-2 mRNA is significantly more stable than iso-1 (Supplementary Fig. 8a), consistent with its prevalence in glioma. Modulation of miR-10b by its inhibitor and mimic had reciprocal effects on the levels of two isoforms, with iso-1 being downregulated and iso-2 upregulated by miR-10b and resulting in the altered total levels of CDC42 protein (Fig. 5c, d).
and Supplementary Fig. 8b, c). Notably, the ASO complementary to 3′ end of U6 snRNA (ASO1) and highly similar to miR-10b, but not another U6 ASO (ASO2), modulated CDC42 splicing and overall expression, similarly to the miR-10b inhibitor (Fig. 5c, d and Supplementary Fig. 8b, c).

miR-10b inhibition reduces glioma growth and leads to glioma cell death in vitro and in animal models in vivo [6, 7, 36]. CDC42 small Rho GTPase has also been implicated in cancer cell proliferation and viability [37]. We, therefore, questioned whether CDC42 could partly mediate miR-10b effects on glioma growth. Downregulation of CDC42 by siRNAs either cognate to the iso-2 or the common region of the iso-1 and iso-2 transcripts (but not by iso-1 specific siRNAs) to the levels mimicking the effect of the miR-10b inhibitor (Fig. 5e and Supplementary Fig. 9a-f), significantly reduced glioma proliferation and growth, as indicated by Ki67 staining, number and size of GSC spheroids, and overall cell viability (Fig. 5f-h and Supplementary
Fig. 9a-f). Conversely, overexpression of CDC42 iso-2 promoted glioma growth similarly to miR-10b (Supplementary Fig. 10a-e). We, therefore, concluded that miR-10b exerts its glioma-promoting activity, in part, by binding U6 snRNA and thereby affecting CDC42 splicing (Fig. 5i). This, in turn, leads to the expression of the more stable CDC42 mRNA isoform and increased protein levels of this important growth-promoting factor.

**Discussion**

MiR-10b has been established as an onco-miR associated with various malignancies [1, 2]. It is most strongly upregulated in malignant gliomas and required for glioma and GSC viability [6, 7]. Nevertheless, there is still little understanding of the striking glioma “addiction” to miR-10b and its signaling in cancer more generally. Even for a well-defined malignancy such as the GBM, the attempts to study molecular mechanisms underlying miR-10b function are hampered by the (a) inter-_and intratumoral heterogeneity of the tumors and multiple mechanisms of miR-10b activity in genetically diverse glioma cells; and (b) unconventional targeting properties of this miRNA [7]. Previous studies suggested that miR-10b commonly binds and regulates target mRNAs by a non-canonical, non-seed-mediated mechanism, and control pre-mRNA alternative splicing [7]. In this study we employed a recently developed CLEAR-CLIP technology for direct transcriptome-wide capture of miR-10b interactions and identification of its targets in glioma. Since the technique is based on the cross-linking and ligation of the spatially proximal transcripts, it may reveal both direct and more complex indirect interactions between the transcripts, including false-positives [14]. Initially proposed quality metrics for CLEAR-CLIP relied on the bioinformatical analysis demonstrating that miRNA-mRNA chimeras are enriched in a miRNA seed. However, such approach is unconstructive for miRNAs that operate via non-canonical binding. Therefore, although our analysis identified more than a hundred of putative targets with miR-10b binding sites (mostly seedless, see below), additional experimental work is required for their functional validation. Here we used several complementary approaches to validate an unanticipated top miR-10b target- a short regulatory snRNA U6, rather than a conventional mRNA. Despite the significant basepairing and abundant binding to miR-10b, U6 was not predicted by current algorithms developed with the focus on mRNA and further updated to include long non-protein-coding RNA species. miR-10b-U6 interactions have been confirmed biochemically, with chimeras observed in AGO2, pan-AGO, as well as snRNP (SART3, PRPF8) immunoprecipitates. Enrichment of miR-10b in high-density nuclear fractions resolved in glycerol gradient, containing pre-catalytic spliceosome, further supports these results. Furthermore, small RNA FISH that we developed for miR-10b detection revealed its co-localization with U6 snRNA and U6 snRNPs in glioma cells and patient-derived tumors. Finally, the chimeric reads produced by miR-10b and U2 snRNA, another spliceosomal RNA and U6-binding partner, further substantiated miR-10b interaction with the spliceosomal machinery.

U snRNAs are essential components of the spliceosome, required for the recognition of substrate pre-mRNAs and serving as ribozyme catalysts of two consecutive transesterification reactions to ligate two exons, concurrent with an intron removal [38]. Cryo-EM structures demonstrated that U6 snRNA forms the catalytic core of the spliceosome and interacts with three other snRNAs (U2, U4, and U5), > 25 protein partners, and the pre-mRNA substrate throughout the splicing cycle, which is accompanied by its large conformational changes [30, 38]. MiR-10b binds to a highly conserved and functionally important part of the U6 at its 3′ end and is anticipated to interfere with the formation of major U6 structures, including internal stem loop (ISL), telostem, and asymmetric bulge (a SART3 binding site) (Fig. 4a). These secondary structures undergo dynamic transitions into other conformations during the spliceosome assembly and activation [39]. For example, incorporation of U6 into the precataytic spliceosomal B complex (one of the key steps of the spliceosome assembly) requires unwinding of the ISL and subsequent annealing to U4 snRNA [30]. The protein SART3 is essential for this reaction [27, 30]. As we demonstrate, it can be displaced from U6 by miR-10b. miR-10b also interacts with PRPF8, a scaffolding factor that mediates the orderly assembly of proteins and snRNAs in the pre-catalytic U4/U6-U5 snRNP complex. The U6 telostem, another miR-10b-interacting structure, is mutually exclusive with the formation of U2/U6 helix II required for the spliceosome activation [40]. Therefore, since U6 secondary and tertiary structures formed in this region are critically important for the spliceosome function and miR-10b modulates these structures, it has a major impact on this fundamental cellular machinery, miR-10b binding may also interfere with essential nucleotides in the U6, such as the highly conserved catalytic AGC triad that is critical for splicing catalysis, and the adjacent ACAGAGA box that interacts with the 5′ splice site of pre-mRNA and helps organize the spliceosome active site through the formation of base triple interactions (Fig. 4a) [41].

MiR-10b also promotes U6 post-transcriptional modifications, including pseudouridylation and m^6A modification. m^6A, the most prevalent internal modification
associated with eukaryotic RNAs, influences many steps of mRNA metabolism and may be important for regulatory functions of ncRNAs as well. In U6 snRNA, a single m6A at position A43 (U6–A43) is modified by a recently identified methyltransferase, METTL16 [42]. METTL16 activity on U6–A43 depends on the formation of the adjacent ISL secondary structure within U6 [43]. While m6A-containing mRNAs undergo distinct pathways of rapid degradation [44], whether and how m6A regulates U6 function and stability in human cells is unknown. Interestingly, the recent yeast-based study demonstrated global splicing alterations in a strain lacking U6–A43 modification and suggested a function for this modification in 5'splice site recognition and exon diversification [45]. This is consistent with the A43 positioned in the center of the ACAGAGA motif essential for U6 base pairing to the intron adjacent to the 5' splice site of a pre-mRNA [41].

Human U6 is also constitutively pseudouridylated at several positions [46], including U6-β that could directly participate in the base-pairing with miR-10b, and U31 and U40, positioned in the telestem and SART3-binding region; they could also be affected by miR-10b and miR-10b-modulated SART3 binding. The effects of these modifications on U6 structure, stability, and functions are largely unknown, although studies of Saccharomyces cerevisiae suggest their functionality [47]. Pseudouridylation is generally thought to be structurally stabilizing because of the high base stacking [48]; however, this notion stems from mRNA studies and may be irrelevant for the highly structured small RNA such as U6. Our study demonstrates that miR-10b increases the levels of U6 pseudouridylation and m6A modification, probably by facilitating the corresponding enzymatic activities, or interfering with the modification-erasing enzymes. Alternatively, miR-10b may modulate these modifications indirectly by affecting U6 conformation and binding to spliceosomal proteins such as SART3. In the context of miR-10b impacting the overall U6 stability, our data suggest the negative effects of the modifications on U6 stability. This could be due to the highly structured character of U6 and binding to multiple RBPs, the characteristics affected by the modifications.

Altogether, our data suggest the mechanistic relationships between miR-10b binding to U6 and its effects on U6 modifications, conformation, and destabilization. However, the molecular mechanisms and the sequence of events remain to be further investigated. We propose that miR-10b binding leads to a U6 structural-conformational reorganization that, in turn, alters protein binding of some snRNP proteins (e.g., SART3, PRPF8) and enzymes catalyzing U6 modification, including writers and erasers (e.g., methyltransferase METTL16 and pseudouridine synthase Pus1). This, in turn, regulates U6 modifications, spliceosome assembly, and splicing reactions. In such a scenario, the lowered levels of U6 could be a consequence of the reduced U6 incorporation in the snRNPs. Alternatively, miR-10b binding may first facilitate the recruitment of U6 modifiers leading to the increased levels of modifications and thereby affecting the U6 structure, steady-state, and splicing cycle. miR-10b binding at the 3'end of U6 can also directly interfere with U6 trimming and turnover. Regardless of the order of events, miR-10b-mediated effects on U6 secondary structure, which is tightly controlled during the splicing cycle, must play a critical role in the downstream effects on mRNA splicing. The data indicating that miR-10b inhibitor and U6 ASO1 that cause similar conformation changes in U6, but reciprocal effects on U6 levels, have nearly identical effects on the CDC42 splicing (and splicing patterns of numerous other genes), further support the importance of miR-10b-mediated U6 reorganization, rather than absolute U6 levels, in splicing catalysis. In part, the structural alterations may result from miR-10b displacement of major U6 binding proteins, such as SART3. Intriguingly, in addition to displacing SART3 from U6, miR-10b also directly reduces SART3 expression by targeting its mRNA [7]. Such a two-fold effect on SART3 may provide a mechanism of enhanced control over U6 structure and function.

One of the important questions regarding miR-10b regulation of U6 snRNA relates to the stoichiometry of the two molecules. U6 is an abundant transcript, which level reaches 100,000 copies per cell in HeLa cells. Many miRNAs also accumulate to very high steady-state levels, with some at least as plentiful as the U6 snRNA [49]. Our analysis of large glioma RNAseq datasets [50], albeit semi-quantitative, suggested that miR-10b levels in glioma have been previously underestimated and may reach the levels of U6. Further optimization of the RNAseq technologies and single-cell analysis will help validate this quantification. In addition, the experiments with Actinomycin D indicate that miR-10b is about 10-times more stable than U6. Furthermore, since it is by far more potent inhibitor of RNA Pol II that transcribes miR-10b (active at 0.5 μg/ml) than of Pol III transcribing U6 (5 μg/ml) [51], our experiments likely underestimate the difference in the stability of the two molecules. The highly stable miR-10b, therefore, could potentially bind to and regulate U6 via a catalytic mechanism, with a single miR-10b copy capable of interfering with multiple U6 RNPs. Since U6 transcript is incorporated in various RNPs that, as recently proposed, could be partitioned in different liquid phases or “droplets” [52, 53], miR-10b may target some of these phases. Furthermore, as mRNA binding to its targets is one of the factors defining the miRNA
half-life [54], ample binding to U6 and U6 RNPs may contribute to the miR-10b steady-state levels.

Our work contributes to the growing body of evidence supporting miRNA nuclear localization and non-canonical functions and expands the mechanisms and repertoire of miRNA targets. In contrast to a canonical pre-miRNA undergoing nuclear export, cleavage by the RNase III Dicer, and loading of a mature miRNA to the RNA-induced silencing complex (RISC) in the cytosol, where it targets mRNAs, some mature miRNAs are found in the nucleus. Among the first reported nuclear miRNAs, for example, is miR-21- another important onco-miR implicated in the carcinogenesis [55]. We demonstrate that about 50% of miR-10b in glioma cells is nuclear, but the subcellular trafficking and selectivity of such localization remain to be further investigated. miR-10b could be either produced by an atypical, entirely nuclear biogenetic pathway or undergo the cytosolic maturation followed by the reverse cytosol-to-nucleus shuttling. Nuclear localization of miRNAs can be mediated by different sequence motifs (e.g., AGUGUU or UUGCAUAGU), in a mechanism controlled by specific RBPs. However, miR-10b does not contain such nuclear localization motifs. At least two pieces of evidence provide support for the nuclear miR-10b maturation scenario: first, the finding of Ago2 and, more recently, of Dicer in the nucleus of specific cells, including glioma (Supplementary Fig. 5a) and [12, 56–58], and second, the co-IP of the spliceosomal SART3 and PRPF8 complexes with Ago2, the core endonucleolytic components of RISC that efficiently pools-down the entire miR-10b population from glioma cells. Our data showing the presence of nuclear Ago2 and its interactions with splicing factors is in agreement with prior reports [59–61]. These data support the scenario of aberrant nuclear miRNA maturation in cancer cells and links nuclear miRNA with the regulation of alternative splicing. There is also emerging evidence that RISC components continuously shuttle between the nucleus and cytoplasm [57]. Although prior data indicate that miRNA-Ago complexes target nuclear transcripts, including mRNAs and pre-mRNAs [56–62], here we characterize the first miRNA target that is a small regulatory RNA essential for eukaryotic cells. This study, therefore, expands the miRNA-regulated networks beyond mRNA and lncRNA targets and suggests exciting new avenues for miRNA research.

In addition to discovering lncRNA as a new class of miRNA targets, our work indicates that an established oncomiR-10b targets multiple transcripts in a noncanonical way, in most cases based on the non-seed and non-3’UTR binding. Although the seed sequences confer the strongest mRNA degradation, functional seedless sites have been discovered in multiple biologically important mRNAs [13, 14, 63, 64]. For example, a new type of miRNA-recognition elements found exclusively in CDSs depend on miRNA 3’-end interactions, but not seed pairing. This type of binding blocks translation elongation in an Argonaute-dependent but GW182-independent manner, resulting in reduced protein but not mRNA levels of the targets [65]. This finding is consistent with our results demonstrating that the majority of seedless miR-10b interactions are found in the CDSs. Of note, some developmentally essential miRNAs, such as miR-9, appear to favor seedless interactions [14]. Furthermore, we demonstrate that seedless interactions can interfere with RBP binding and modulate target’s structure, thereby affecting its properties, and regardless of the effects on the target stability. Such miRNA control could be especially important for altering structural characteristics and activity of regulatory transcripts, including small and long ncRNAs. Investigation of the normal physiological and pathology-associated networks formed by interacting miRNAs and other regulatory RNA species represents an exciting new avenue for fundamental and translational research.

Conclusions

This study demonstrates a new and fundamental layer of miRNA regulation in cancer, extending it beyond mRNA targeting. It presents an unexpected intersection of miRNA and splicing machineries and highlights the importance of unbiased investigation of molecular mechanisms mediated by regulatory RNAs in cancer.

Abbreviations

GSC: Glioma stem cells; CLEAR-CLIP: Covalent ligation of endogenous Argonaute-bound RNAs-crosslinking and immunoprecipitation; SnRNA: Small nuclear RNA; EGF: Epidermal growth factor; FGF: Fibroblast growth factor; RT-qPCR: Quantitative Reverse Transcription –PCR; FISH: Fluorescence In-situ hybridization; ISL: Internal stem loop.

Supplementary information

The online version contains supplementary material available at https://doi.org/10.1186/s12943-022-01494-z.

Additional file 1 Supplementary Fig. 1. Analysis of putative miR-10b binding sites in 5’UTR, CDS, and 3’UTR of mRNA targets identified by CLEAR-CLIP, using a STARmiRDB tool [66], demonstrates the distribution of seed-based and seedless targets. Related to Fig. 1.

Additional file 2 Supplementary Fig. 2. Relative levels of miR-10b in glioma and non-glioma cell lines. The qRT-PCR reactions were performed with equal RNA input and the Ct values are indicated.

Additional file 3 Supplementary Fig. 3. Additional data for miR-10b and U6 FISH in cells and tissues. a Representative FISH images of pri-miR-10b (red) and U6 (green) in cultured LN229 and U251 glioma cells with the corresponding fluorescently labeled probes, and nuclei stained with DAPI. b Representative FISH images of the miR-10b paralogue miR-10a (red) and U6 (green) in patient-derived GBM tissues demonstrate low miR-10a expression and lack of colocalization. c Representative FISH images of miR-10b (red) and U6 (green) in human kidney tissues.
El Fatimy et al. Molecular Cancer (2022) 21:17

Additional file 4 Supplementary Fig. 4. MiR-10b is enriched in the spliceosome in glioma cells. a Western blot analysis of LN229 and U251 nuclear fractions separated in glycerol gradient, demonstrates enrichment of the spliceosomal marker PRP8 in the heavy fractions. b qRT-PCR analysis demonstrates miR-10b enrichment (C1 values < 25.5) in the corresponding heavy fractions in both cell lines.

Additional file 5 Supplementary Fig. 5. SART3 and PRP8 bind to AGO proteins in glioma cells. a Representative images of miR-10b FISH (red) and AGO2 immunofluorescence (green) in glioma cells, and nuclei stained with DAPI (blue). Arrows mark the colocalization of AGO2 and miR-10b. b-e SART3 and PRP8 bind to AGO proteins in glioma cells. IP with either IgG (control), AGO2, SART3, or PRP8 antibodies, followed by the Western blotting for the indicated proteins.

Additional file 6 Supplementary Fig. 6. MiR-10b regulates U6 snRNA but not U2 and U4 snRNAs in glioma cells. a Representative denaturing Northern blotting of glioma cells and GSCs transfected with either U6 ASOs, miR-10b inhibitor or mimic, with probes specific for U2 and U4 snRNAs. b MiR-10b mimic and inhibitor do not regulate 3′ UTR luciferase reporters bearing full-length U6 snRNA sequence in glioma cells. The data is presented as Renilla/Firefly relative luminescence and normalized to the corresponding values in cells not transfected with the oligonucleotides. n = 12; Graphical data are shown as mean ± SEM. c qRT-PCR analysis of snRNA U6 levels in U251 cells transfected with either miR-10b inhibitor, mimic, or corresponding controls, and treated with 5 μg/ml Actinomycin D (mean ± SD, n=3). d U251 cells were treated with Actinomycin D followed by the qRT-PCR analysis of miR-10b and U6 levels (mean ± SD, n=3). e Levels were calculated using two-way ANOVA. f U251 cells were treated with Actinomycin D followed by Western blot analysis at the indicated time points (mean ± SD, n=3). P values were calculated using two-tail unpaired t-test. *P < 0.05; ***P < 0.001.

Additional file 7 Supplementary Fig. 7. Relative levels of U6 snRNA in normal neuroglial and glioma cells. a qRT-PCR analysis of U6 levels in normal human neuroglial cells (neuroprogenitors (NSC), neurons, and astrocytes) and glioma cells. The expression of U6 was normalized by uniformly expressed miR-125a. b qRT-PCR analysis of U6 levels in human astrocytes transfected with miR-10b mimic, or corresponding control oligonucleotides (mean ± SD, n=3). c Levels were calculated using two-tail unpaired t-test. **P < 0.01.

Additional file 8 Supplementary Fig. 8. Effects of miR-10b and U6 snRNA on CDC42 alternative splicing. a Differential stability of CDC42 iso-1 and iso-2 variants. LN229 cells have been treated with Actinomycin D followed by qRT-PCR analysis of CDC42 iso-1 and CDC42 iso-2 variants. b qRT-PCR analysis of CDC42 iso-1 or CDC42 iso-2 levels in GBM8 cells transfected with either U6 ASOs, miR-10b inhibitor, mimic, or corresponding control oligonucleotides (mean ± SD, n=3). Fold-change in the expression of the isoforms is plotted relative to the corresponding control conditions. P values were calculated using two-tail unpaired t-test. c Western blotting analysis of the indicated proteins in GBM8 cells transfected with either U6 ASOs, miR-10b inhibitor, or mimic. *P < 0.05; **P < 0.01.

Additional file 9 Supplementary Fig. 9. CDC42 promotes the growth of GSCs. a GBM8 cells were transfected with siCDC42 iso-2, siCDC42 total, or control siRNAs, followed by the qRT-PCR analysis. b GBM8 cells were transfected with siCDC42 iso-2, siCDC42 total, or control siRNAs, followed by Western blotting analysis. KD of the CDC42 iso-2 reduced total CDC42 levels. c The growth of GSC spheroids, untreated or transfected with CDC42-targeting siRNAs, have been analyzed. The number and size of GSC neurospheres have been calculated at day 7 after transfections (mean ± SD, n=3). P values were calculated using two-tail unpaired t-test. *P < 0.05; **P < 0.01; ***P < 0.001; ns, no significance.

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Authors’ contributions
AMK and REL conceived and designed the study; REL and YZ performed experiments; data analysis, and visualization; ED and ZW contributed to data analysis; NMT contributed to CDC42 splicing analysis; MR, HS, RR, and EJU assisted with experiments; REL, YZ, and AMK wrote the manuscript. All authors revised and approved the manuscript.

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Availability of data and materials
All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this
paper may be requested from the authors. The RNA-seq data are deposited to GEO with accession number GSE182280. The CLEAR-CLIP data are deposited to GEO with accession number GSE182281.

Declarations

Ethics approval and consent to participate

De-identified human brain tumor specimens were obtained from the DFCI/BWH Brain Tumor BioRepository, according to protocol approved by the Institutional Review Board.

Consent for publication

All authors have agreed to publish this manuscript.

Competing interests

All other authors declare no competing financial interests.

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