Supplementary Information for

Somatic LINE-1 Promoter Acquisition Drives Oncogenic FOXR2 Activation in Pediatric Brain Tumor

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Abbreviations for Fig. 1b:
CNS tumors of 17 entities, Abbreviations: CHL/AD: child/adult; DIA/DIGG: Desmoplastic infantile astrocytoma/gangliogioma; EPN: Ependymoma; CNS NBL FOXR2: FOXR2-activated CNS neuroblastoma; G3: Group 3; G34R: Histone H3 G34R-mutant diffuse hemispheric glioma; G4: Group 4; GBM: Glioblastoma; IHG: Infant-type hemispheric glioma; INF: Infant; K27M: Histone H3 K27M-mutant diffuse midline glioma; MB: Medulloblastoma; MES: Subclass mesenchymal; MID: Subclass midline; MYCN: Subclass MYCN; RTKI/II: Subclass RTKI/II; SHH: SHH-activated medulloblastoma; WNT: WNT-activated; YAP: YAP1-fused; ZFTA: ZFTA-fused

Additional Results and Discussion:

Histopathological and molecular characterization of the index tumor samples
The second recurrent tumor sample of our index patient was subjected to our clinical cancer genomic profiling by three-platform sequencing of whole-genome, whole-exome, and
transcriptome [31], and harbored two additional somatic pathogenic variants: a homozygous clonal TP53 R175H mutation and a subclonal PDGFR A amplification (Supplementary Fig. 3). The patient was initially diagnosed with HGG at 11 months of age, with a tumor that showed both low- and high-grade histology and a wild-type expression pattern of p53 as demonstrated by immunohistochemistry (Supplementary Fig. 3e). After receiving chemotherapy treatment, the high-grade glioma relapsed a year later and contained a clonal TP53 mutation, a common feature of pediatric HGG but exceptionally rare for CNS NBL FOXR2. After irradiation treatment, a year later, the tumor recurred for a second time with an even stronger expression of mutant p53 (Supplementary Fig. 3e) and a subclonal copy gain of PDGFR A identified by methylation array (Supplementary Fig. 3c), both are common in pediatric HGG but not CNS NBL FOXR2. The subclonal PDGFR A amplification, potentially associated with conferring therapeutic resistance, was observed only in the second recurrent tumor. These findings were corroborated by both WGS and FISH (Supplementary Fig. 3d). Interestingly, the DNA methylation profiles of the primary and recurrent tumors (1st and 2nd) all clustered with reference tumors of CNS FOXR2-altered NBL, suggesting a common lineage related to FOXR2 activation (Fig. 1b). The results of DKFZ methylation classifier (molecularneuropathology.org) also support this finding (calibrated scores 0.23804, 0.72338, and 0.99994 for CNS NBL FOXR2 for the primary and 1st and 2nd recurrent tumors, respectively). We were able to perform additional whole-exome and RNA-seq on the primary tumor sample acquired at diagnosis. Exome data showed the absence of any pathogenic coding variants at diagnosis, consistent with the wild-type TP53 status by immunohistochemistry. More importantly, RNA-seq data displayed transcription of the non-canonical FOXR2 isoform as well as the L1/FOXR2 fusion transcript (Supplementary Fig. 2a),
confirming the presence of the somatic L1 insertion and activation of *FOXR2* expression driven from the inserted L1 promoter sequence at diagnosis.

**Prevalence of somatic L1 insertion in pediatric high-grade gliomas**

While *de novo* somatic L1 insertions have been studied extensively in adult cancers [16, 33, 34, 38], and epithelial cancers show higher L1 mobilization and subsequent *de novo* insertion rates as compared to brain and blood cancers [1, 20], yet the activity levels of these elements in pediatric cancers have not been well studied. Following the discovery made on the index HGG, we performed a screening of somatic L1 insertions on 87 HGG paired tumor/normal WGS data sets by filtering somatic insertions identified by the structural variant caller Manta [4] and by running the mobile element detector MELT [14] and TraFiC-mem [29](Supplementary Methods). We identified three additional pediatric tumor samples, each showing evidence of a single somatic L1 insertion, albeit intergenic events (Supplementary Methods, Supplementary Table 3), consistent with the low L1 insertion rate observed in CNS tumors previously [1, 20].

**Prevalence of L1 promoter donation in pediatric brain tumors**

To determine the prevalence of transcript activation by this L1 “promoter donation” mechanism, we analyzed tumor RNA-seq from an additional 182 pediatric HGG samples and 22 previously published CNS tumors in a study that reported *FOXR2*-activated CNS neuroblastoma as a new brain tumor entity [36]. We screened for the presence of fusion transcripts involving splicing of an L1 97 SD sequence to a genomic location. While we identified several transcripts matching this profile, most were likely caused ‘spuriously’ by nearby known polymorphic L1s or older L1 lineage insertions as has been previously described [2], or exhibited characteristics of ‘exonization’[30, 32, 39] with additional upstream reads also displaying splicing to L1 sequence (Supplementary Table 2 and Supplementary Fig. 2c).
Promoter donation as a new L1-mediated oncogenic mechanism
Repurposing an active promoter of a retrotransposed L1 for oncogene activation is a new
mechanism in human cancers unveiled by the in-depth genomic, transcriptomic, and epigenomic
analysis presented in this study. The transduction sequence observed in the FOXR2 L1 somatic
insertion indicates the event was most likely a retrotransposition event of the 6p24.1 L1 which
was identified as one of the top five “hot” L1 source elements that gave rise to half of all somatic
L1 insertion events in adult cancers in a recent pan-cancer study [29]. The transduction occurred
with an L1 mRNA transcript that bypassed the canonical L1 polyadenylation signal (5’-
AATAAA) and terminated at an alternative downstream polyadenylation signal (5’-AATATA).
Furthermore, the presence of the poly-A tail, TSDs and an endonuclease (EN)-cut site (5’-
TCTTT/AT) indicates that the FOXR2 L1 somatic insertion event occurred by canonical L1 EN-
mediated retrotransposition.

In contrast, previously reported driver somatic L1 insertion events in adult cancers revealed
mutagenesis by disruption, i) either disruption of a tumor suppressor gene (i.e., APC in
colorectal cancer [25, 34] or PTEN in uterine cancer [16]), or ii) disruption of a repressive
regulatory element allowing expression of an oncogene (i.e., ST18 in liver cancer [35]), or iii)
creation of an intronic novel cis-enhancer driving expression of a putative oncogene (i.e., STCI
in ovarian cancer [27]). Although a previous pan-cancer analysis of transposable elements cited
several examples of oncogene overexpression accompanied by nearby somatic L1 insertions, it
should be noted that all these nearby L1 insertions were 5’ promoter-truncated (e.g., lacking
5’UTRs), thus ineligible for promoter “donation” [29]. Importantly, through the analysis of
serial samples, we show that the somatic L1 insertion was the initiation event present in the
primary tumor, while the other pathogenic variants, e.g., TP53 mutation and PDGFRA amplification, were detected only at tumor recurrence. Thus, promoter “donation” by somatic L1 retrotransposition represents a new mechanism for oncogenesis in human cancers and potentially for aberrant activation of genes involved in other human diseases. It should be noted that popular somatic L1 insertion detection tools, such as MELT [14] and TraFiC-mem [29], the latter of which was previously used for large-scale pan-cancer analysis of WGS data, were unable to identify the somatic FOXR2 L1 insertion identified in this study [14, 29], suggesting that these events may be underrepresented in the current somatic L1 insertion datasets and may be contributing to more pathogenic alterations than currently appreciated.

Materials and Methods:

Data sets
All WGS and RNA-seq data were downloaded from the St. Jude Cloud (www.stjude.cloud) [24]. WGS were mapped to GRCh38 (GRCh38_no_alt) with BWA-MEM [22] alignment of RNA-seq were mapped to GRCh38 (GRCh38_no_alt) by STAR [9]. Specifically, the index tumor is a 2nd recurrent high grade glioma (SJHGG030242_D1) which, along with its matching normal (SJHGG030242_G1), was DNA and RNA extracted and profiled by the three-platform WGS, exome, and transcriptome sequencing employed by the Genome for Kids study and patient consent was given as previously stated [26]. HGG RNA-seq (n=182) and paired tumor-normal WGS (n=87) used for recurrence screening was profiled by the Pediatric Cancer Genome Project [11], Genome for Kids [26], or Real-Time Clinical Genomics (Supplementary Table 5). We also downloaded and analyzed the RNA-seq data of 22 previously published CNS tumors (EGAD00001001927) which were used for identifying a new brain tumor entity, FOXR2-
activated CNS neuroblastoma [36]. Additionally, the authors of the publication provided us with
FPKM values from Affymetrix array FOXR2-activated CNS neuroblastomas (GEO accession:
GSE73038) [36]. We downloaded and processed 453 GTex V7 normal brain tissue samples (99
Anterior cingulate cortex (BA24), 130 Cortex, 117 Frontal Cortex (BA9), and 107
Hippocampus) and compared the expression of FOXR2 with FPKM per gene calculated in
RSEM [21] based on Gencode v31 annotation. The L1 consensus sequence used for analysis
comparisons was that of L1.3 described and published previously [10].

RNA isolation and sequencing on SJHGG030242 primary tumor
A PureLink FFPE Total RNA Isolation Kit (Thermo Fisher Scientific) was used for total RNA
extraction from formalin-fixed paraffin-embedded (FFPE) tumor tissue as previously described
[6, 8, 15]. Purified RNA was quantified on a Qubit 3 Fluorometer (Thermo Fisher Scientific)
using Qubit™ RNA BR Assay Kit (Thermo Fisher Scientific). Total RNA sequencing was
performed using the Illumina TruSeq Stranded Total RNA protocol with at least 500 ng of total
RNA. The quality of the starting materials was checked with the RNA 6000 Nano Assay on a
2100 Bioanalyzer (Agilent) or the RNA Pico Sensitivity Assay on a LabChip GX Touch
(PerkinElmer). Libraries were prepared using the TruSeq Stranded Total RNA Sample Prep Kit
(Illumina), followed by library quantification through qPCR using Quant-iT™ PicoGreen
dsDNA Assay Kits (Thermo Fisher Scientific) or KAPA Library Quantification Kits for Illumina
platforms (KAPA Biosystems), and through low pass sequencing on a MiSeq Nano v2
(Illumina). All sequencing data were generated after 100 cycles of paired-end runs on an
Illumina HiSeq 2500 or HiSeq 4000.

DNA isolation and whole exome sequencing on SJHGG030242 primary tumor
Genomic DNA was extracted from formalin-fixed paraffin-embedded (FFPE) tumor tissue using a QIAamp DNA FFPE Tissue Kit (Qiagen) as previously described [15]. At least 250 ng of genomic DNA was used for each sample. DNA quality was assessed on a 4200 TapeStation (Agilent). Genomic DNA libraries were generated using the SureSelectXT Kit (Agilent Technologies), followed by exon enrichment using the SureSelectXT Human All Exon V7 bait set (Agilent Technologies). The resulting exon-enriched libraries were subjected to paired-end, 100-cycle sequencing performed on a HiSeq 4000 (Illumina).

Methylation array analysis
Genomic DNA extracted from formalin-fixed paraffin-embedded (FFPE) tumor samples was used for genome-wide methylation profiling and CNV analysis by the Illumina Infinium MethylationEPIC platform as previously described [5-8, 15, 17, 23]. At least 250 ng of genomic DNA was used for each sample. For comparison, publicly available well-characterized reference methylation profiles of brain tumors were downloaded from the Genomic Data Commons Data Portal (https://portal.gdc.cancer.gov/). Analysis of methylation profiles, including normalization, filtering, t-distributed stochastic neighbor embedding (t-SNE), and CNV analysis was performed in R v4.1.0 using ChAMP v2.22.0 [37], minfi v1.38.0 [13], limma v3.48.0 [28], conumee v1.26.0, and Rt sne v0.15 (with theta = 0.0) packages in Bioconductor v3.13 (http://bioconductor.org/) as previously described [5-8, 15, 17, 23]. Raw signal intensities were normalized by performing background correction and a dye-bias correction for both color channels with the functional normalization method. The following filtering criteria were applied: removal of probes targeting the X and Y chromosomes; removal of probes containing single-nucleotide polymorphisms; and removal of probes not mapping uniquely to the human reference
genome (hg19), allowing for one mismatch, after removal of poor-quality (P > 0.01) and failed probes. Beta values of the 5000 most variable CpG sites were derived for t-SNE analysis.

**Histopathology review and immunohistochemistry**
Histopathology was centrally reviewed by a neuropathologist specializing in pediatric CNS tumors (J.C.). Standard hematoxylin and eosin histopathologic preparations from each tumor were supplemented by immunohistochemistry on 5-μm formalin-fixed, paraffin-embedded (FFPE) tissue sections. Monoclonal anti-p53 antibody (Zeta Corp, Z2029M, clone DO-7, diluted 1:200) was used to identify the nuclear accumulation of mutant p53. Staining was performed on a Ventana Benchmark Ultra automated stainer with 20-minute CC1 pretreatment and 32-minute incubation with the antibody at 37 °C. IVIEW DAB detection kit was used to visualize the staining results.

**Interphase fluorescence in situ hybridization**
Amplification of *PDGFRA* (4q12) was detected by interphase fluorescence in situ hybridization in a Clinical Laboratory Improvement Amendments (CLIA)-certified laboratory in St. Jude Children’s Research Hospital with probes developed in-house using the following BAC clones: *PDGFRA* (RP11-231C18 + RP11-601H15) with 4p12 control (CTD-2057N12 + CTD-2588A19), as previously described [15].

**Targeted Amplification and PacBio Sequencing**
Targeted amplification of the region on chromosome X containing the putative L1 insertion was performed using extracted DNA from the 2nd recurrent tumor (SJHGG030242_D1) and paired normal (SJHGG030242_G1) with Kapa HiFi HotStart Ready Mix KR3070, 300 nM final concentration of the flanking primers (Line_ChrX-For/Line_ChrX-Rev), and 25 ng of genomic
DNA in a 25 ul reaction volume. After the initial denaturation at 95°C for 5 minutes, PCR was completed with 35 cycles of amplification consisting of a 20 second denaturation at 98°C, 15 second annealing at 62°C, and an 8 minute extension at 72°C. A final extension of 10 minutes was performed prior to a 4°C hold step. A ~3,800 bp PCR fragment amplified from the 2nd recurrent tumor which contains the putative LINE-1 insertion and a 423 bp fragment from the matching paired normal (SJHGG030242_G1) were used to prepare the PacBio sequencing libraries following the “Procedure & Checklist-Preparing SMRTbell® Libraries using PacBio® Barcoded Overhang Adaptors for Multiplexing Amplicons”. The libraries were loaded at 6 pM and run on the PacBio Sequel system using 1 1M v3 LR SMRT Cell.

The PacBio generated circular consensus sequence (CCS) fastq files were then visually analyzed and compared to the L1.3 consensus sequence with SerialCloner v2.6.1 to determine the details of the inserted sequence. Sequence quality generated by PacBio using CCS reads is of high quality based on previously published data showing the ability of PacBio to successfully sequence through L1 poly-A sequence [12].

Bisulfite Conversion, PCR amplification, and Illumina MiSeq Sequencing
Non-FFPE tissue DNA samples in this study were bisulfite converted using the Zymo EZ DNA Methylation-Gold Kit D5006, with 500 ng of DNA under standard protocol conditions.
Amplification of the chr6p24.1 source element was performed using the previously published primers in Tubio et al., 6p24.1 F (chr6:13190940-13190966; hg19) and 6p24.1 R (L1Hs:206-229)(Supplementary Table 4, chr6_tubio_for_bs_conv and chr6_tubio_rev_bs_conv) [38]. The 25ul PCR reaction tube (Micron Low bind 0.2 ml PCR tube) contained Kapa HiFi HotStart Uracil + Ready Mix KK2801, Gibco Ultra Pure H2O (DNase/RNase free), 300 nM each
chr6_tubio_for_bs_conv and chr6_tubio_rev_bs_conv PCR primers. All PCR reactions were performed in a BioRad C1000 thermocycler with an initial denaturation for 3 minutes at 95°C, followed by 38 cycles of a 20 second denaturation at 98°C, a 15 second 57°C annealing, and a 30 second 72°C extension with a final extension of 5 minutes at 72°C, followed by a 4°C hold. Note that primer designs for the 6p24.1 source element excluded the last 6 CpG sites captured in the Xp11.21 FOXR2 insertion described below.

Bisulfite converted chrX-L1 DNA was amplified with the same above PCR reagents, except with a mixture of the following primers (600 nM ChrX-L1_bs_full-d2-For; 300 nM each ChrX-L1_bs_methyl-d2-Rev, ChrX-L1_vs_conv-site1-Rev, ChrX-L1_bs_conv-site2-Rev, ChrX-L1_bs_both-sites-Rev). The ChrX primers were designed to overlap the ORF2/5’UTR junction sequence of the FOXR2 L1 insertion. The thermocycling settings for amplification included an initial denaturation at 95°C for 3 minutes, followed by 38 cycles of 98°C denaturation for 20 seconds, 55°C annealing for 15 seconds, and 72°C extension for 30 seconds, and a final extension at 72°C for 1 minute, followed by a 4°C hold.

After PCR, primers and nucleotides were removed using a 1:1 ratio of Beckman Coulter Ampure beads A63880. Libraries were generated from the amplicons using the Roche Kapa Hyperprep DNA library kit. The amplicon library (4-6 nM) was loaded onto an Illumina MiSeq. Sequencing was performed using a 500-cycle v2 Nano reagent kit operating with v4 control software, and local run manager v3.0

Analysis of Bisulfite Converted Illumina MiSeq Fastq Reads
The generated 250 bp pair-end Miseq reads were trimmed using trimgalore v0.4.4 and cutadapt v1.8.1 with the default parameters to remove the adaptors.

For analysis of bisulfite sequence reads of the L1 5’UTR of the chr6p24.1 source element, trimmed MiSeq fastq reads were aligned with bismark v0.23.1dev [19] to the expected flanking genomic location and the 5’UTR of the chr6 amplified source element sequence [14](Suppl. Table S9D; element 6:13191033), requiring that all aligned reads align with no indels present. Passed reads were analyzed for presence of bisulfite converted nucleotides.

For analysis of the ORF2/5’UTR chrX FOXR2 L1 sequence, the trimmed 250 bp paired-end reads were first merged into single-end reads based on overlap with bbmerge (bbmap, version 38.86, sourceforge.net/projects/bbmap/) set with parameter “forcemerge” to true [3]. Single-end reads ranging in size from 350 to 400 bp, the expected length of the inverted ORF2/5’UTR amplicon were aligned with bismark v0.23.1dev [19] to the expected sequence, requiring no indels in analyzed reads, and resulting reads displayed hypo-methylation.

**Analysis of FOXR2 Splicing Pattern in the Index HGG RNA-seq Samples**

To confirm that FOXR2 expression in the index HGG was caused by L1 promoter “donation” instead of L1 exonization, we generated a template sequence for the L1-inserted genomic region by inserting the ~3kb PacBio sequence outlined in Figure 1d to the L1 insertion site at chrX: 55,597,061 (GRCh38). We then map all RNA-seq reads to this L1-inserted template sequence by running STAR v2.7.9a in two-pass mode.

Reads aligned exclusively within the inserted L1 were removed as they could represent the expression of nearly identical L1 located in other genomic regions. For the same reason, reads
mapped within the repetitive regions of the template sequence were removed. Reads with >90% identity to the template sequence were retained for examining the splicing pattern. The resulting RNA-seq read alignments are presented in Supplementary Fig. 2a using the IGV viewer. Somatic insertion of L1 5’ UTR being the transcription initiation site can be verified by the following two patterns: 1) there is no splicing between the region upstream of the L1 insertion and the inserted cryptic L1, which excludes the possibility of L1 exonization; 2) all splice junction reads connecting to the acceptor site of FOXR2 exon 2 were from the L1 5’ UTR 97 donor sites; which shows that transcription only occurred on the L1-inserted mutant allele.

**Detecting L1 Fusion Transcript Involving 97 Donor Site in Pediatric Brain Tumors RNA-seq**

We used the L1.3 consensus sequence as the “bait” to identify chimeric reads with partial match to L1. Mapped RNA-seq bam files downloaded from the St. Jude Cloud were converted to fastq using Picard v2.6.0 and subsequently trimmed using trimgalore v0.6.7 and cutadapt v3.1 with the default parameters to remove the adaptors. The fastq files downloaded from EGAD0001001927 from the 22 previously published CNS tumors were trimmed as above [36] and underwent the same analysis as further. The trimmed fastq files were then mapped to the L1.3 consensus sequence [10] using bowtie2 v2.2.9 with “--very-sensitive-local parameter”. Reads with edit distance \( \leq 2 \), a threshold set to ensure we only retain reads likely belong to the young L1 elements similar to L1.3, were kept for future analysis.

To identify aberrant splicing events involving the L1 97 splice donor (SD) site, similar to the index HGG, we developed a custom script to extract the reads with a minimum overlap of 20 bp to the 97 SD, i.e. 78-97 bp of L1.3 in the remapped L1.3 bam file. For reads that pass this threshold and contain soft-clipped (SC) subsequences unaligned to L1.3, the soft-clipped reads
were then extracted and aligned to GRCh38 (GRCh38_no alt) to find the target exons by running bowtie2 v2.2.9 using the same parameters as above. Those that have ≥2 high-confidence match (<2 mismatches) to the reference genome were retained and annotated on the presence of L1 elements within 20 kb based on annotated polymorphic L1 on the Repeats Track: RepeatMasker of the UCSC genome browser (http://genome.ucsc.edu) [18] or somatic or germline L1 insertion identified by MELT v2.2.2 or Manta (details below).

For the 22 previously published CNS tumors we only assessed for presence of L1/FOXR2 fusion transcripts and unfortunately did not identify any and thus did not include these samples in Supplementary Table 2 nor Supplementary Table 3 results.

Identification of somatic L1 insertions using paired tumor-normal WGS data set
Manta v1.6.0 [4] was run with default settings on the paired WGS data and somatic insertions were manually inspected for bi-directional soft-clipped reads representing L1-related subsequence and poly-A tail. We confirmed that HGG WGS of our index sample can be detected by this approach. When using this approach to analyze the entire cohort, we identified a second event in a different HGG in an intergenic region on chromosome 7 (Supplementary Table 3).

MELT v2.2.2 [14] was run, first running the preprocess mode and then Melt-single mode only to L1 sequences as described at (https://melt.igs.umaryland.edu/manual.php) on tumor and matched normal samples independently. Events with a “PASS” filter were retained and further assessed manually for soft-clipped read support in IGV bam viewer for the presence of somatic L1 integration events. MELT did not identify the L1 insertion site in the index sample; however, it did discover two additional L1 insertion sites in our cohort (Supplementary Table 3).
TraFiC-mem was run through the docker container with default settings on the paired tumor-normal WGS data of our index case (GRCh37 BWA aligned) (https://gitlab.com/mobilegenomesgroup/TraFiC)[29] and no somatic L1 integration sites were identified. The results were also negative in the additional 3 HGG tumors, which had somatic L1 insertions detected by MELT and Manta (Supplementary Table 3) as well as the rest of the cohort with paired tumor-normal samples. GRCh37 BWA alignments were used for analysis as TraFiC-mem can only run on hg19 aligned samples.

Supplementary Fig. 1. L1 Insertions in Index Patient a Schematic of non-canonical FOXR2 isoform (blue) on the X chromosome (red vertical bar) with a cluster of soft-clipped reads (SC,
red arrow) observed in tumor whole genome sequencing below. Black arrows mark the forward (F) and reverse (R) primers used for PCR amplification results in supplementary Fig. 1b shown. b Amplification of the genomic region flanking the SC cluster at chrX:55597061 (GRCh38). The HGG displays a ~3.8kb amplicon (red arrow) while the matching germline sample a 423bp band as expected from the reference genome. Thermofisher’s 1kb Plus ladder is shown on left. c WGS evidence of the 6p24.1 L1 source element in the tumor (top) and normal (bottom) with support of poly-A tail (green box) and L1 5’UTR (gray box).
Supplementary Fig. 2. RNA Splicing Patterns

a Splicing pattern in HGG RNA-seq supporting FOXR2 activation via L1 promoter donation. Top, a schematic drawing of the promoter donation model where the inserted L1 serves as exon 1 for transcription initiation. Bottom, mapping of RNA-seq reads from the primary and 2nd recurrent HGG to a template sequence representing the mutant allele with somatic L1 insertion in the non-canonical intron 1 of FOXR2. The inserted cryptic L1 sequence which was determined by targeted PacBio sequencing shown in
Supplementary Table 1 is labeled as L1Hs_INSERTION. The absence of splicing from the genomic sequence upstream of the L1 insertion, which includes the non-canonical exon 1, to either the inserted L1 or the non-canonical exon 2 supports transcription initiation at L1 5’ UTR (only first 97bp of L1 5’UTR are labeled). b Unique transcript reads supporting the splicing of L1 97 splice donor to FOXR2 splice acceptor in the primary and 2nd recurrent tumor transcriptome. c An example of L1 exonization based on screening of 182 HGG RNA-seq. Top, schematic drawing of L1 exonization which involves splicing of an upstream exon to the inserted L1 site. Bottom, a L1 exonization example identified in RNA-seq data of SJHGG030665_D1 in the LINC00907 region where L1PA4 is integrated at intron 7 of LINC00907 as part of the reference human genome. RNA splicing was detected in the genomic region upstream of the L1PA4 integration site (i.e., exon 7 and intron 7 in L1ME4A) to 5’UTR of L1PA4 in addition to splicing from L1PA4 to the downstream exon 8 of LINC00907, which matches the pattern of the L1 exonization model. d Unique transcript reads supporting the exonization example described in Supplementary Fig. 2c. Periods represent the additional genomic sequence present between the junctions shown.
Supplementary Fig. 3. Molecular and genetic data of the HGG progression samples from the index patient. 

a) Summary table of molecular and genetic analysis done on respective samples.
samples (N.T. means not tested). b Summary table of genetic variants discovered in the HGG progression samples. c Chromosomal copy number variations of the primary diagnosis sample (top), 1st recurrent tumor (middle), and 2nd recurrent tumor (bottom) analyzed by methylation array. A focal PDGFRA amplification was detected only at the 2nd recurrent tumor. d Fluorescence in situ hybridization (Red: PDGFRA, 4q12; Green: Control, 4p12) in the primary tumor (left) and 2nd recurrent tumor sample (right). White arrows indicate cells with PDGFRA amplification detected only at the 2nd recurrent tumor. d p53 IHC stains are shown for respective tumors.
**Supplementary Table 1.**
Sequence of flanking amplified *FOXR2* locus including the L1 insertion sequence and the subsequent matching SNPs to chr6p24.1 source element.

**Supplementary Table 2.**
Results of L1 97 splice donor transcript screening in 182 pediatric high grade glioma tumors.

**Supplementary Table 3.**
Additional intergenic somatic L1 insertions identified by Manta and MELT in the pediatric HGG cohort.

**Supplementary Table 4.**
Primer names and sequences used and described in materials and methods.

**Supplementary Table 5.**
A list of the pediatric high-grade glioma diagnosed sample ids, age of diagnosis, associated cohort, and type of sequencing data analyzed from the St. Jude Cloud.
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