Detection of a Distinctive Genomic Signature in Rhabdoid Glioblastoma, A Rare Disease Entity Identified by Whole Exome Sequencing and Whole Transcriptome Sequencing

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Abstract
We analyzed the genome of a rhabdoid glioblastoma (R-GBM) tumor, a very rare variant of GBM. A surgical specimen of R-GBM from a 20-year-old woman was analyzed using whole exome sequencing (WES), whole transcriptome sequencing (WTS), single nucleotide polymorphism array, and array comparative genomic hybridization. The status of gene expression in R-GBM tissue was compared with that of normal brain tissue and conventional GBM tumor tissue. We identified 23 somatic non-synonymous small nucleotide variants with WES. We identified the \textit{BRAF} V600E mutation and possible functional changes in the mutated genes, \textit{ISL1} and \textit{NDRG2}. Copy number alteration analysis revealed gains of chromosomes 3, 7, and 9. We found loss of heterozygosity and focal homozygous deletion on 9q21, which includes \textit{CDKN2A} and \textit{CDKN2B}. In addition, WTS revealed that \textit{CDK6, MET, EZH2, EGFR,} and \textit{NOTCH1,} which are located on chromosomes 7 and 9, were over-expressed, whereas \textit{CDKN2A/2B} were minimally expressed. Fusion gene analysis showed 14 candidate genes that may be functionally involved in R-GBM, including \textit{TWIST2} and \textit{UPK3BL}. The \textit{BRAF} V600E mutation, \textit{CDKN2A/2B} deletion, and \textit{EGFR/MET} copy number gain were observed. These simultaneous alterations are very rarely found in GBM. Moreover, the \textit{NDRG2} mutation was first identified in this study as it has never been reported in GBM. We observed a unique genomic signature in R-GBM compared to conventional GBM, which may provide insight regarding R-GBM as a distinct disease entity among the larger group of GBMs.
Introduction
Rhabdoid glioblastoma (R-GBM) is a very rare disease with few cases reported [1–8]. R-GBM is characterized by tumor cells that resemble rhabdomyoblasts [2], which robustly express vimentin, epithelial membrane antigen (EMA), and SMARCB1 (INI-1), but only faintly express glial fibrillary acidic protein (GFAP) [5,7,9,10]. Clinically, R-GBMs can occur at any age but most commonly occur in teenagers younger than 20 years old [1,2,5].

Chromosome 22, which is frequently lost in atypical teratoid rhabdoid tumors (ATRT), is often deleted in these tumors [3,8], although this finding is inconsistent [1]. In one case, copy number gains were noted for chromosomes 3, 7, 9, 12, 17q, and 21q [1] in R-GBM. Also, in a case series, copy number gain or amplification of EGFR on chromosome 7 was noted [9]. Regarding genetic changes prevalent in brain tumors, CDKN2A hemizygous deletion was reported in one case [2]. Otherwise, BRAF mutations were absent in two cases that were examined [11], and SMARCB1 (INI-1) [12,13], which is important in ATRT, was not mutated in R-GBM [2].

Presently, R-GBM is not recognized as a distinct disease entity by the World Health Organization [14] classification system because accumulated information on this rare variety is still rudimentary. To our knowledge, no study has evaluated the genome-wide profile of this disease except for one case that was evaluated using array comparative genomic hybridization (CGH) [1]. To determine whether R-GBM should be recognized as a disease that is distinct from conventional glioblastoma (GBM) or other tumors with similar characteristics such as ATRT, comprehensive genomic data will be fundamental for diagnostic, prognostic, and therapeutic decisions.

Materials and Methods

Study Patient
A 20-year-old female patient was seen in an outpatient clinic at Seoul National University Hospital because of headache, nausea, and vomiting in April 2011. Brain magnetic resonance imaging showed a 5-cm sized, well-enhanced mass in the right temporal lobe. The mass also showed diffusion restriction with increased perfusion at the peripheral enhanced portion. She underwent a craniotomy for tumor removal in May 2011. The molecular genetic characteristics of the surgical specimen were evaluated as follows. Immunohistochemical staining revealed focal expression of GFAP and strong expression of EMA and INI-1 (Figure 1). Fluorescence in situ hybridization (FISH) showed no EGFR amplification and no deletion of chromosomes 1p, 9p21, or 19q. In addition, methylation-specific PCR showed hypermethylation of the MGMT promoter, and the MIB-1 labeling index was measured as 36.5% with an Aperio Spectrum plus image analyzer. The study patient received adjuvant concurrent extensive surgeries and concurrent chemoradiotherapy combined with oral temozolomide treatment. She remained free of disease for 25 months after the treatment. Using next generation sequencing techniques, we studied this tumor to obtain novel insight into identifying distinctive genetic changes in an R-GBM compared to conventional GBM as well as normal brain tissue. We performed whole exome sequencing (WES), whole transcriptome sequencing (WTS), single nucleotide polymorphism (SNP) array, and array-CGH. The aims of this study were to investigate the genomic profile of R-GBM and to explore whether R-GBM had a distinct genomic signature that could be used as a therapeutic target.

![Figure 1. Pathology of rhabdoid glioblastoma.](image-url)
and quality were assessed using an Agilent 2100 Bioanalyzer (Agilent Grand Island, NY) and eluted in RNAse-free water. RNA quantity RNA was extracted from the tumor tissue using TRIzol (Invitrogen, Extracted DNA was quantified using a NanoDrop ND-1000 logy Information (http://www.ncbi.nlm.nih.gov/) to assist in locating clusters were downloaded from the National Center for Biotechno-
dedFuse [19], BreakFusion [20], and ChimeraScan [21]. UniGene unknown bases or low quality bases, the remaining reads were aligned to the reference genome, mapping and pairing were performed with the Burrows-Wheeler Aligner (BWA) algorithm[15]. Local realignment was performed using Genome Analysis ToolKit (GATK) [16], and duplication removal was conducted using Picard.

Somatic calling of somatic single nucleotide variants (SNVs) and indels is described in Supplement 1. Using SnpEff [17], we selected variations that were non-synonymous and rare in the general population (defined as <1% in the 1000 genome project (http://www.1000genomes.org/)). For copy number alteration (CNA) analysis of WES data, we used the Copy Number Analysis for Targeted Resequencing (CONTRA) tool [18] and summarized the exon-level log2 fold changes of read depth between the normal and tumor samples into gene-level log2 fold changes. Loss of heterozygosity (LOH: heterozygous in normal tissue but homozygous in the tumor) analysis also was performed using WES. We used variant allele fraction values of normal and tumor samples to determine the LOH region.

Whole Exome Sequencing
We used the Agilent SureSelect50-Mb ExomeCapture Kit for exon target enrichment (Agilent Technologies Inc.). Sequencing was performed using the Illumina HiSeq2000 (Illumina Inc., San Diego, CA) with 100-bp paired-end reads. Using UCSC hg19 as a reference genome, mapping and pairing were performed with the Burrows-Wheeler Aligner (BWA) algorithm [15]. Local realignment was performed using Genome Analysis ToolKit (GATK) [16], and duplication removal was conducted using Picard.

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Whole Transcriptome Sequencing
The 200- to 500-bp double-stranded cDNA fragments were purified by agarose gel electrophoresis and amplified using PCR to produce the library. Raw sequencing reads were produced by Illumina HiSeq 2000 with 100-bp paired-end reads. After removing noisy raw reads, which contained the adaptor sequence and more than 10% unknown bases or low quality bases, the remaining reads were aligned with the human reference genome (UCSC hg19). To find fusion transcripts, we utilized three types of fusion discovery software: deFuse [19], BreakFusion [20], and ChimeraScan [21]. UniGene clusters were downloaded from the National Center for Biotechnol-
y and quality were assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies Inc., Santa Clara, CA).

DNA and RNA Preparation
Fresh frozen tumor tissue and 5 ml peripheral blood were obtained at the time of the first surgery. The DNeasy® Blood & Tissue Kit (Qiagen, Hilden, Germany) was used to extract genomic DNA and tumor DNA, according to the manufacturer’s recommendations. Extracted DNA was quantified using a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE). RNA was extracted from the tumor tissue using TRIzol (Invitrogen, Grand Island, NY) and eluted in RNAse-free water. RNA quantity and quality were assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies Inc., Santa Clara, CA).

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SNP Array
We applied a genome-wide SNP array (Illumina HumanOmmi5-Quad BeadChip, Illumina) using the genomic DNA sample. With B allele frequency data from GenomeStudio (Illumina) analysis results for SNP array data, we used the paired parent-specific circular binary segmentation method for LOH and CNA analysis.

Array Comparative Genomic Hybridization and Identification ofCNAs
We used the Agilent aCGH G3 Human 1×1M array with tumor and matched normal genomic DNA samples. Raw data were acquired and normalized using the locally weighted scatterplot smoothing (LOW-ESS) algorithm using Feature Extraction software ver10.7 (Agilent software). The significance test for each CNV region used the Z-statistic calculated by DNA Analytics ver4.0.81 (Agilent software), which sets the window size to 1M and Z-score threshold to 4.0.

Use of the Public Database as a Reference
We used gene expression data estimated from WTS to select possible functional genetic changes in our study. Because R-GBM is a rare disease and obtaining control samples is not easy, we used a public database as a reference. First, we compared the RPKM value of specific genetic changes found in our analysis with normal brain expression values. Then, we compared the RPKM value of specific genetic changes found in our analysis with GBM data to determine whether R-GBM is simply a subtype of GBM. For the normal brain data, we used the normalized expression dataset from BrainSpan (http://www.brainspan.org/). For the GBM data, we used datasets from TCGA (https://tcga-data.nci.nih.gov) and cBioPortal for Cancer Genomics (http://www.cbioportal.org).

Results
Tumor Purity, Alignment, and Coverage Statistics
The purity of the tumor samples was estimated using SNP array data with the Allele-specific copy number analysis of tumors (ASCAT) algorithm [24]. The proportion and the ploidy of tumor cells in the sample were about 89% and 2.17, respectively (Online Resource Section 1: Supplementary Figure 1). In WES, the total numbers of uniquely mapped reads were 181,350,341 and 186,695,100 for normal and tumor samples, respectively. These data yielded mean target coverages of 210 and 197 for the samples, respectively (Online Resource Section 2: Supplementary Table 1).

Somatic SNVs and Small Indels Found With WES
We found 46,468 (45,045 in dbSNP138) and 46,191 (44,748 in dbSNP138) SNVs from the paired normal DNA and tumor DNA, respectively. A total of 45,542 (44,264 in dbSNP138) SNVs were identified in both samples (Online Resource Section 2: Supplementary Table 1). We also identified 3753 (3362 in dbSNP138) small indels from the paired normal DNA samples and 3678 (3314) small indels from the paired normal DNA samples.

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detected with WES. Twenty-three non-synonymous SNVs (Table 1) were found, 13 of which were also found with WTS.

Loss of Function SNVs and Analysis of Small Indels (Online Resource Section 5)

The candidates for loss of function were selected from the nonsense, splice junction, and frameshift variants (Supplementary Table 3 of Online Resource Section 5).

Whole Chromosome Copy Gains and Losses

Gains were identified in chromosomes 3, 7, and 9 from the SNP array and WES data (Figure 2). Interestingly, chromosome 9 showed a homozygous deletion of the 9p21 locus that contains the tumor suppressor genes CDKN2A and CDKN2B (Figure 2B).

CNA Analysis Using WES and Array-CGH

CNA data were generated from array-CGH and WES. First, CNA were analyzed with array-CGH with probe-based CNA and interval-based CNA. The probe-based method revealed that more than 80,000 CNA regions were present in tumor tissue compared to paired normal tissue. The interval-based method revealed 370 tumor-specific CNA regions. WES analysis identified 323 regions with CNAs, which included 11 genes that are well-known tumor suppressors and oncogenes [25,26]: VHL, CTNNB1, PIK3CA, EGER, CDK6, MET, EZH2, MLL3, CDKN2A, CDKN2B, and NOTCH1 (Table 2). In addition, WES, SNP array, and array-CGH analyses showed that CDKN2A/2B were homozygously deleted. Copy number gain was observed for CTNNB1, CDK6, VHL, MLL3, EZH2, PIK3CA, EGER, NOTCH1, and MET with WES and SNP array analyses.

Fusions Found With WTS

A total of 376 fusions were observed with WES with deFuse [19], BreakFusion [20], and ChimeraScan [21]. Intrachromosomal fusions and intrachromosomal fusions >50 kb were selected, and thus, 24 fusions (Online Resource Section 6: Supplementary Table 4) were analyzed further. Among these 24 fusions, in-frame fusions were selected for candidate genetic hallmarks in R-GBM.

Selection of Genetic Hallmarks in R-GBM

We used WTS data to investigate functional genetic changes in R-GBM, and the public database was used as a reference. First, we compared the RPKM values of specific genetic changes found between tumor and normal brain tissue (Figure 3). We focused on affected genes with more than a 4-fold change in expression and integrated the results among WES, WTS, and array-CGH. Several genes had significant SNVs, CNAs, or fusions. Among genes with SNVs, NDRG2, NKA1N2, CER1, and ISL1 were downregulated, whereas PARP9 was upregulated in the tumor sample of the study patient compared to normal brain tissue. Among genes with CNAs, NOTCH1, EGER, CDK6, EZH2, and MET were upregulated, whereas CDKN2A and 2B were downregulated in the tumor sample of the study patient compared to normal brain tissue. These results are summarized in Table 3.

The aforementioned analysis was assumed to have identified functional genetic changes in the selected genes. In addition, RPKM values of these selected genes in GBM and R-GBM were compared (genetic changes in GBM were obtained from the TCGA database). We listed genes with more than a 2-fold difference in genetic expression between conventional GBM and R-GBM. The following significant alterations between study samples (R-GBM) and conventional GBMs were found: 1) CER1 and ISL1 had SNVs that were significantly downregulated. 2) CDKN2A and 2B were genes with CNAs and were significantly downregulated. 3) NOTCH1, EGER, CDK6, PIK3CA, and MET were genes with CNAs and were significantly upregulated. 4) PD1, RASSF8, FBKP15, GALNT6, ITGA6, SLC6A6, TWIST2, and UPK3BL were significantly up-regulated fusion genes correlated to R-GBM.

Search for GBMs With Similar Genetic Hallmarks in the TCGA Database

In our case, genetic hallmarks excluding fusions are summarized below (Table 3): 1) BRAF V600E; 2) NDRG2 192F and ISL1 C234W mutation; 3) CDKN2A/2B homozygous loss; and 4) EGER, CDK6, EZH2, NOTCH1, and MET copy number gain. Subsequently, we searched the cBiportal (TCGA provisional data) for

Table 1. List of 23 Candidate Non-Synonymous Somatic SNVs

| Chr | Position | dbSNP | Ref | Alt | Transcript | Gene | Effect | AA Change | Depth o | Depth v | VAF o | VAF v | RNA-Seq (confirmed) |
|-----|----------|-------|-----|-----|------------|------|--------|-----------|---------|---------|--------|--------|---------------------|
| Ch1 | 27876290 | . | G | C | NM_001029882.2 | ADHC1 | MISSENSE | H779Q | 81 | 66 | 0 | 0.409 | O |
| Ch1 | 104093621 | . | G | C | NM_0176103.3 | BNP3 | MISSENSE | P474A | 169 | 192 | 0 | 0.432 | O |
| Ch2 | 27403455 | . | G | A | NM_182591.3 | LCLAT1 | NONSENSE | W42* | 266 | 259 | 0 | 0.409 | X |
| Ch2 | 209201623 | . | G | A | NM_015040.3 | PIKK3YE | MISSENSE | G152R | 182 | 174 | 0 | 0.379 | O |
| Ch3 | 52413954 | . | G | A | NM_151512.4 | DAIH1 | MISSENSE | E2471K | 15 | 34 | 0 | 0.294 | O |
| Ch3 | 122247474 | . | T | C | NM_031458.2 | PARP9 | MISSENSE | T768A | 234 | 300 | 0 | 0.563 | O |
| Ch5 | 50685703 | . | C | G | NM_002202.2 | ISL1 | MISSENSE | C234W | 78 | 77 | 0 | 0.455 | X |
| Ch6 | 12469435 | . | G | A | NM_00104251.1 | NKN2 | NONSENSE | V471 | 278 | 264 | 0 | 0.371 | X |
| Ch7 | 42962956 | . | C | T | NM_002247.9 | PSM2A | MISSENSE | G142R | 169 | 176 | 0 | 0.477 | O |
| Ch7 | 99170311 | . | A | T | NM_001083956.1 | ZNF655 | MISSENSE | M229L | 183 | 204 | 0 | 0.309 | O |
| Ch7 | 140453136 | n11348022 | A | T | NM_004334.3 | BR4 | MISSENSE | V600E | 208 | 294 | 0 | 0.551 | O |
| Ch8 | 14720162 | . | G | A | NM_005454.2 | CER1 | MISSENSE | P211S | 163 | 183 | 0 | 0.006 | 0.874 | X |
| Ch10 | 100185398 | . | C | G | NM_000195.3 | HP51 | MISSENSE | S203T | 64 | 63 | 0 | 0.397 | O |
| Ch11 | 799344 | . | G | A | NM_145886.3 | PIDD | MISSENSE | S899F | 53 | 67 | 0 | 0.373 | O |
| Ch11 | 124766783 | . | G | A | NM_019055.5 | ROBO4 | MISSENSE | R119W | 25 | 23 | 0 | 0.478 | X |
| Ch11 | 111186936 | . | A | G | NM_00103403.2 | ZCCHC16 | MISSENSE | I138N | 65 | 141 | 0 | 0.142 | X |

Chr, Chromosome; Ref, Reference; Alt, Alternative; AA, amino acid; VAF, Variant allele frequency.
GBM cases that also harbor the above mutations. For mutations, we used WES data (284 samples), and for gene copy number gain/loss, we used array-CGH data (497 samples).

Among 284 GBM cases with sequencing data, the \textit{BRAF} V600E mutation was found in 1.7% of GBM cases. The \textit{NDRG2} mutation was not found in GBM. However, when we searched for \textit{NDRG2} genetic changes in the array-CGH database, \textit{NDRG2} was amplified in two cases \((2/497 = 0.4\%\)) and homozygously deleted in three cases \((3/497 = 0.6\%\)). For \textit{ISL1}, no mutation was found in GBM cases. However, the \textit{ISL1} deletion was observed in two GBM cases \((2/497 = 0.4\%\)).

Based on array-CGH data from 497 GBM cases, homozygous deletions of \textit{CDKN2A} and \textit{CDKN2B} were found in 62\% and 61\% of

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Copy number status of R-GBM. (A) Gross copy number changes, (B) variant allele frequency in the tumor sample, and (C) variant allele frequency in the normal sample.}
\end{figure}
Copy number gains of \( \text{EGFR} \), \( \text{CDK6} \), \( \text{EZH2} \), \( \text{NOTCH1} \), and \( \text{MET} \) were found in 49%, 7%, 4.4%, 0.6%, and 8.9% of the cases, respectively. Interestingly, \( \text{NOTCH1} \) copy number gain, and that of \( \text{EGFR} \), \( \text{CDK6} \), \( \text{EZH2} \), and \( \text{MET} \), were mutually exclusive. Two out of three \( \text{NOTCH1} \)-amplified cases accompanied the \( \text{CDKN2A/B} \) homozygous deletion. On the other hand, copy number gains in \( \text{EGFR} \), \( \text{CDK6} \), \( \text{EZH2} \), and \( \text{MET} \) were not mutually exclusive, and co-amplification of these genes was frequently seen.

In three GBM cases with the \( \text{NDRG2} \) homozygous deletion, two had simultaneous homozygous \( \text{CDKN2A} \) and \( \text{CDKN2B} \) deletions.

### Table 2. Genes with Copy Number Alterations that Are Well Known to be Associated with Cancer Development and/or Progression

| Chr | Start       | End         | Gene    | Genetic alteration | aCGH | WES | SNP array | Classification | Expression Ratio1 * | Expression Ratio2 # |
|-----|-------------|-------------|---------|--------------------|------|-----|----------|----------------|---------------------|---------------------|
| Chr3| 9022276     | 9291369     | \( \text{VHL} \) | Gain               | Yes  | Yes | Yes      | TSG            | -0.28               | 3.07                |
| Chr3| 41240942    | 41281939    | \( \text{CTNNBI} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | -0.04               | 5.83                |
| Chr3| 178916609   | 17892393    | \( \text{PIK3CA} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 1.93                | 2.92                |
| Chr7| 55080951    | 55214485    | \( \text{EGFR} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 3.02                | 1.54                |
| Chr7| 92234235    | 92465941    | \( \text{CDK6} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 3.05                | 1.86                |
| Chr7| 116312459   | 116438440   | \( \text{MET} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 4.32                | 3.83                |
| Chr7| 148504464   | 148581441   | \( \text{EZH2} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 5.29                | 0.12                |
| Chr7| 151832010   | 152133090   | \( \text{MLL3} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 0.84                | 1.94                |
| Chr9| 21967751    | 21994490    | \( \text{CDKN2A} \) | Loss               | Yes  | Yes | Yes      | TSG            | -2.38               | -1.03               |
| Chr9| 22005935    | 22009013    | \( \text{CDKN2B} \) | Loss               | Yes  | Yes | Yes      | TSG            | -5.02               | -8.89               |
| Chr9| 13938896    | 139440238   | \( \text{NOTCH1} \) | Gain               | Yes  | Yes | Yes      | Oncogene       | 2.37                | 2.77                |

Chr, chromosome; aCGH, array comparative genomic hybridization; WES, whole exome sequencing; Del, deletion; TSG, tumor suppressor gene.

* Expression ratio 1 is the log2 ratio of the expression level (value in RPKM) in our patient over the mean expression level (value in RPKM) in normal brain (http://www.brainspan.org).

* Expression ratio 2 is the log2 ratio of the expression level (value in RPKM) in our patient over the mean expression level (value in RPKM) in glioblastoma multiforme (https://tcga-data.nci.nih.gov/tcga/tcgaHome2.jsp).

Figure 3. Expression status of selected genetic changes found in the R-GBM sample in comparison with normal brain. (A) genetic changes with copy number alteration, (B) genetic changes with single nucleotide variation, and (C) genetic changes with gene fusion (y axis shows the log2 ratio of the expression level (value in RPKM) in our patient over the mean expression level (value in RPKM) in normal brain (http://www.brainspan.org)).
amplification at the same time. The BRAF mutation, MET, EGFR, and CDK6 amplifications, and CDKN2A/B homozygous deletion at the same time. However, only three among 497 cases had NOTCH1 amplification at the same time. The BRAF mutation and EZH2 amplification were also mutually exclusive. To summarize, although our case of R-GBM is not representative of all R-GBMs, coexistence of the genetic hallmarks found in our patient is a very rare event in GBM.

**Discussion**

In this study, we addressed the genomic profile of R-GBM, a very rare disease entity. At the chromosomal level, we found copy number gains in chromosomes 3, 7, and 9, and the deletion of 9p21. When we correlate this karyotypic abnormality with genetic changes, we made the following observations.

On chromosome 3, PIK3CA was amplified, and its corresponding expression was elevated compared to normal brain. PIK3CA is frequently altered in GBM, and indeed, amplification of this gene is found in 13% of primary GBMs [27]. Hence, the PIK3CA copy number gain found in our sample was not surprising and implies that a common genetic denominator exists between GBM and R-GBM.

On chromosome 7, EGFR, EZH2, CDK6, and MET had copy number gain, and their expression was elevated compared to that in normal brain tissue. In fact, the gain of chromosome 7 along with EGFR and MET gene amplification is relatively common in adult brain tumors including GBM [28]. In addition, EGFR copy number gain and amplification were observed in a series of R-GBM cases [9,29]. On the other hand, amplification of EZH2 and CDK6 is not commonly observed; only 4% and 7% of GBM cases had amplification of these genes, respectively. Moreover, 1.2% of GBM cases had co-amplification of EZH2 and CDK6, and 0.4% (2 out of 497) of GBM cases also had co-amplification of MET, EZH2, and CDK6 according to the TCGA database. We reviewed the pathology slides of these two cases, which are detailed on the websites (cBioportal case id=TCGA-06-0187 and cBioportal case ID=TCGA-19-1390). A pathology review of these two cases did not provide a definite diagnostic clue regarding R-GBM. Therefore, we could not draw a definite conclusion regarding whether amplification of one or more of EZH2, CDK6, or MET may be an irrelevant event or an oncogenic driver in the pathogenesis of R-GBM.

On chromosome 9, NOTCH1 copy number gain and associated over-expression were observed. Although the role of the Notch pathway in brain tumors is an area of active investigation, Notch1 signaling is known to promote survival of GBM cells via EGFR-mediated signaling [30]. In addition, Notch signaling has oncogenic potential in a model of medulloblastoma [31]. Hence, we believe that the NOTCH1 copy number gain found in our case may have substantially contributed to oncogenesis and tumor progression. However, only three among 497 cases had NOTCH1 amplification in the TCGA database, which implies that this alteration in NOTCH1 is not a common event in GBM. Regarding the genomic profile of these three cases, two harbored the CDKN2A/2B homozygous deletion as in our case, and one case harbored TP53 and IDH1 missense mutations. As mentioned in the Results section, NOTCH1 amplification was mutually exclusive with EGFR, MET, EZH2, and CDK6 amplification in the TCGA database. Hence, the simultaneous copy number gain in, and over-expression of, NOTCH1, EGFR, MET, EZH2, and CDK6 in our sample is a very interesting phenomenon. What is most interesting regarding NOTCH1 amplification is that one case with NOTCH1 amplification in the TCGA database (cBioportal case id=TCGA-02-2483) had rhabdoid features upon pathology review. Therefore, we believe that further testing for NOTCH1 copy number gain in other R-GBM samples is necessary to confirm whether NOTCH1 is a key factor for rhabdoid morphogenesis.

For chromosome 9, the 9p21 deletion (rather than chromosome 9 copy number gain) was found using WES and SNP microarray. This alteration was not detected with conventional FISH, which confirms the high sensitivity of WES and SNP microarray compared to conventional FISH. Chromosome 9p21 contains CDKN2A and CDKN2B, which are well-known tumor suppressor genes that play an important role in GBM. CDKN2A and CDKN2B were homozygously deleted in this patient, and their expression was correspondingly low. Thus, CDKN2A and CDKN2B may play an important role in our patient.

As for non-synonymous SNVs excluding BRAF V600E, we designated the ISL1 and NDRG2 mutations as genetic hallmarks of R-GBM. We selected these genes for the following reasons. First, gene expression of ISL1 and NDRG2 was significantly reduced compared to expression in normal brain, which implies that these genetic changes are functional. Second, both ISL1 and NDRG2 are biologically relevant to brain tumor development. ISL1 is required for neural development, and expression of this gene is associated with neuroendocrine carcinoma [32,33]. NDRG2 is a well-known tumor suppressor in brain tumors [34]. In contrast to our sample, ISL1 and...
was identified in a small subset (0.4% and 0.6%, respectively) of NDRG2 (n = 3) in the GBM (TCGA database), oncogenes including MET amplifications were found in two cases (67%). Amplification of other NDRG2 were found in which was recently shown to be important in ATRT \[37\], was observed factor for rhabdoid morphogenesis. However, \[12,35,36\], were not altered in R-GBM as had previously been shown.

Finally, comparison of genetic changes in our case with those of ATRT is valuable because ATRT and R-GBM share common morphologic features. First, genes in the SWItch/Sucrose NonFermentable (SWI/SNF) complex, which is a genetic hallmark of ATRT \[12,35,36\], were not altered in R-GBM as had previously been shown. This finding suggests that although ATRT and R-GBM share common morphologic features, the SWI/SNF complex abnormality is not a key factor for rhabdoid morphogenesis. However, EZH2 over-expression, which was recently shown to be important in ATRT \[37\], was observed in our sample. EZH2 was both amplified and over-expressed in our sample. Therefore, EZH2 copy number gain and over-expression may play an important role in rhabdoid tumor generation.

Here, we addressed genetic hallmarks found in our R-GBM case including BRAF V600E, ISL1 C234W, NDRG2 192F, CDKN2A/2B deletion, NOTCH1 copy number gain, and gain of chromosome 7 (including CDK6, MET, EZH2, and EGFR copy number gain). The patterns of mutation and gene expression in R-GBM are rather unique compared to conventional GBM, suggesting that R-GBM is a distinct disease entity. Among these genetic changes, NOTCH1 copy number gain and NDRG2 mutation, which are rare events in the TCGA GBM database, appear to be important genetic markers in R-GBM formation. Furthermore, EZH2 copy number gain and over-expression may play an important role in rhabdoid tumorigenesis.

Appendix A. Supplementary data
Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.tranon.2015.05.003.

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