Somatic mutations in myeloid cell leukemia-1 contribute to the pathogenesis of glioma by prolonging its half-life

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Abstract. The identification of mutated genes in glioblastoma multiforme (GBM) is an essential step towards improving current understanding of the molecular mechanism underlying the disease and establishing novel targets for diagnostic and therapeutic purposes. The present study used direct sequencing to screen 20 malignancy-associated genes, which have either been well described in the literature or observed multiple times in human cancer sequencing, in cancerous and normal control tissue samples from 20 patients with histologically confirmed GBM. The investigation identified five somatic non-synonymous coding mutations in four candidate genes, with two located in the proline, glutamic acid, serine, threonine-rich region of myeloid cell leukemia sequence 1 (Mcl-1), (D155G and L174S). The sample pool was then expanded by sequencing Mcl-1 in a further 43 patients with GBM and another somatic mutation in the same region, D155H, was identified. The subsequent functional investigation confirmed that these somatic mutations affected the degradation of Mcl-1, and the growth of glioma cells transfected with mutant plasmids was significantly accelerated compared with cells overexpressing wild-type Mcl-1. The mutational profiling of GBM in the present study revealed for the first time, to the best of our knowledge, several mutations in Mcl-1, and identified this gene as a novel therapeutic target for the treatment of GBM.

Introduction

Glioma is a type of tumor, which grows from glial cells, a supportive cell in the brain, and is the most life-threatening type of brain tumor (1). Glioblastoma multiforme (GBM) is the most advanced and aggressive subtype of glioma, and patients with GBM have a median survival rate of ~14 months (2). As a complex medical condition, GBM is considered to be result from the interaction between multiple genetic and environmental factors (3). Currently, only a small number of low penetrance variants attributing to cancer risk have been identified using genome-wide association investigations (4), and the etiology remains to be elucidated. Previous efforts to comprehensively characterize the genomes of primary GBM have established that the disease is driven by numerous and diverse genetic events in individual patients (5,6).

It is generally accepted that somatic mutations are important in the pathogenesis of cancer, and that cancer develops via the accumulation of somatic mutations in certain cancer-specific genes, including oncogenes and tumor suppressors, depending on the type of tumor (7). Previous studies have demonstrated that the frequency of somatic mutations in candidate cancer genes is significantly higher than expected, and that the combination of certain individual mutations may have a specific effect on the properties of the tumor (8-11). These mutations are considered to result from a combination of environmental and genetic factors (12). Following the determination of the human genome sequence, several investigations have been performed to determine somatic mutations in various types of cancer. The Sanger sequencing technique was used to directly sequence 13,023 genes, identifying 189 genes carrying excessive somatic mutations in human breast and colorectal cancer (10). A mismatch repair detection method was used to screen 22 cell lines and 93 matched tumor-control sample pairs, for somatic mutations in 30 cancer-associated genes A total of 152 somatic mutations were identified in breast and colorectal cancer (13), including genes, which are reported to be involved in the development of cancer, including Kirsten rat sarcoma virus oncogene and v-raf murine sarcoma viral oncogene homolog B (BRAF).

To further investigate the prevalence and distribution of somatic mutations in glioma, the present study aimed to examine 63 glioma tissues and their matched normal tissues for somatic mutations in 20 genes, which have been reported to as affirmatively or potentially associated with oncogenesis (14,15). A total of six somatic mutations were identified, with three repeated somatic mutations observed exclusively in myeloid cell leukemia sequence 1 (Mcl-1). The effect of these somatic mutations were subsequently investigated.

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Materials and methods

Subjects and tumor samples. A total of 63 patients (42 male and 21 female) with histologically confirmed GBM were recruited at the Department of Neurosurgery, The Affiliated Hospital of Peking University (Beijing, China), where the patients received surgical resection. The median age of the patients was 55 years (range, 45-67 years) and none had received any preoperative treatment. All the tumor specimens were examined by two independent, experienced pathologists prior to establishing the final diagnosis. High-fidelity polymerase chain reaction (PCR) and direct sequencing were performed to screen the coding region of 20 genes (Mcl-1, Bcl-2, EGFR, KRAS, FGR, IKZF1, BTG1, TP53, PAX5, BRAF, BACH2, ARF, Bcl6, VEGF, HER2, β-catenin, c-myc, Rh, EZH2, E2F) for potential somatic mutations in 20 patients, and the gene of interest was further screened in the remaining 43 patients. The size of the tumor was determined by measuring the maximum tumor diameter presented on radiographic images, including computed tomography scans and magnetic resonance imaging. The tumor tissue samples were rapidly frozen immediately following surgical resection in liquid nitrogen and stored at -80°C for future investigations. Peripheral blood samples (5 ml) were acquired from all patients. Informed consent was obtained from the patient's families, and the present study was approved by the Review Board of the Hospital Ethics Committee at Dalian Medical University (Dalian, China).

DNA extraction and nucleotide sequencing analysis. Total genomic DNA was isolated from the GBM tumor tissue samples and peripheral blood using a DNA extraction kit (Invitrogen Life Technologies, Carlsbad, CA, USA). The isolated DNA was dissolved in sterilized Milli-Q water (EMD Millipore, Billerica, MA, USA) and quantified using a NanoDrop spectrophotometer (Thermo Fisher Scientific, Wilmington, DE, USA). Agarose electrophoresis (Sigma-Aldrich, St. Louis, MO, USA) was used to confirm the purity of the DNA. The chromosome segments comprising the coding region of each gene were amplified by PCR (primer set: forward 5'-GTATCGAGCTAGCGCTCCGCTATG-3' and reverse 5'-GTACTCTTCAGCGAGCTAGAT-3'; Sangong Biotech, Hangzhou, China) according to the following PCR cycling conditions: 95°C for 10 min; 35 cycles of 95°C for 30 sec, 58°C for 30 sec, 72°C for 1 min, followed by 72°C for 10 min. The PCR products were purified using an ExoSAP-IT purification kit (United States Biochemical Corp., Cleveland, OH, USA), prior to sequencing using an ABI sequencing system by PerkinElmer Applied Biosystems (Foster City, CA, USA). The sequences were subsequently aligned against the revised Cambridge sequence in the MITOMAP database (www.mitomap.org) using the MegAlign program from the DNASTAR software package 12.1 (DNASTAR, Inc., Madison, WI, USA). The sequence alterations identified in the tumor tissue samples, but not in the matched peripheral blood samples, were recorded as somatic mutations. The screened somatic mutations were confirmed at least twice by additional independent PCR and resequencing.

Cell lines and cell culture. The U251 GBM cell line was cultured in RPMI-1640 medium (Invitrogen Life Technologies) containing 10% fetal bovine serum (Sigma-Aldrich Canada Ltd., Oakville, Ontario, Canada), 100 U/ml penicillin and 100 mg/ml streptomycin (Invitrogen Life Technologies).

Apoptotic assay. Apoptosis was determined using an annexin V/fluorescein isothiocyanate (FITC) apoptosis detection kit (Keygen Biotech Co., Ltd., Nanjing, China), according to the manufacturer's instructions. Briefly, glioma cells (2x10^6 cells) cultured in 10 cm dishes were trypsinized (Invitrogen Life Technologies), washed with phosphate buffered saline (PBS) and subsequently stained with FITC-conjugated anti-annexin V antibody in the dark for 15 min at room temperature. The cells were then analyzed by flow cytometry (FACSCanto II; BD Biosciences, San Jose, CA, USA). All experiments were performed in triplicate.

Half-life determination. 35S methionine (Guidechem, Hangzhou, China) pulse-chase labeled glioma cells over-expressing wild-type Mcl-1 (D155G, D155S or L174S) were plated into media deficient in methionine for 1 h at 37°C. The cells were washed with PBS and pulsed with labeling media containing 35S methionine, at a final concentration of 100 mCi/ml, and the synthesis of Mcl-1 was inhibited by exposure to ultraviolet light. The cells were washed and harvested at 0, 0.5, 1 and 2 h. and treated with 1 mg/ml recombinant TRAIL (R&D Systems, Minneapolis, MN, USA) for 30 min. Immunoprecipitation. The cells transfected with wild-type or mutant Mcl-1 were incubated with beads coupled to antibody. Protein A agarose beads (Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA; 0.5 ml bed volume) were incubated in 1 mg/ml BSA in 1 ml phosphate-buffered saline containing 100 μg polyclonal anti-Mcl-1 antibody (Santa Cruz Biotechnology, Inc.) at 4°C overnight. Following incubation, the beads were pelleted by centrifugation and washed three times. The radioabeled cells were harvested and lysed in cold radioimmunoprecipitation (RIPA) lysis buffer, containing 10 mM Tris-HCl (pH 7.4), 1% NP-40, 1 mM EDTA, 0.1% sodium dodecyl sulphate (SDS) and 150 mM NaCl.
supplemented with fresh proteinase inhibitor cocktail (Sigma-Aldrich), followed by breaking through a 25 gauge syringe 20 times. Equivalent quantities of the cell extract were adjusted to equal volumes using RIPA buffer, precleared with protein A agarose beads and clarified by centrifugation at 400 x g for 10 min at 4°C. The lysates were subsequently incubated with anti-Mcl-1 antibody at 4°C overnight and the immune complexes were precipitated using protein A agarose beads for 6 h at 120°C. The immunoprecipitates were washed three times with RIPA buffer and incubated at 95°C for 5 min in SDS sample buffer. The protein samples were separated on an SDS-polyacrylamide gel electrophoresis (PAGE) gel, followed by autoradiography (Biospace Lab, Nesles la Vallée, France) to detect the signals.

Western blot analysis. U251 cells were transfected with wild-type or mutant Mcl-1 and were lysed in lysis buffer (Beyotime, Shanghai, China) on ice 72 h after transfection. The cell lysates were loaded onto 10% SDS-PAGE gels (Invitrogen Life Technologies) and the separated proteins were transferred onto a polyvinylidene fluoride membrane (EMD Millipore), which was subsequently blocked with Tris-buffered saline, containing 5% non-fat dry milk at room temperature for 1 h. The membrane was subsequently incubated with the mouse monoclonal anti-haemagglutinin tag antibody (cat. no. 12ca5; Roche Diagnostics, Mannheim, Germany; 1:1,000) and rabbit polyclonal anti-Mcl-1 antibody (cat. no. sc-819; Santa Cruz Biotechnology, Inc.; 1:2,000) at 4°C overnight, followed by incubation with horseradish peroxidase-anti-rabbit secondary antibody (Cell Signaling Technology, Inc., Danvers, MA, USA; cat. no. 7074S) and goat-anti-mouse (cat. no. sc-2031; 1:10,000; Santa Cruz Biotechnology, Inc.) at room temperature for 1.5 h. Chemical fluorescence was detected using an enhanced chemilluminescence kit (Amersham Biosciences, Piscataway, NJ, USA), according to the manufacturer's instructions. The target bands were densitometrically analyzed and normalized against actin.

Statistical analysis. Statistical analysis was performed using SPSS 19.0 software (SPSS, Inc., Chicago, IL, USA). The
LV et al.: SOMATIC MUTATIONS IN MYELOID CELL LEUKEMIA–1 PROLONG ITS HALF-LIFE

association between somatic mutations and clinicopathologic parameters of GBM was examined using Fisher’s exact test. P<0.05 was considered to indicate a statistically significant difference.

Results

Identification of genetic determinants. To identify the genetic determinants of GBM, the present study initially performed Sanger sequencing of 20 cancer-associated genes in the tumor tissue samples and the corresponding blood genomic DNA from 20 patients with histologically confirmed GBM. Comparison between the tumor and corresponding blood DNA revealed five somatic mutations in the coding regions of four of the 20 candidate genes, with two somatic mutations located in the proline, glutamic acid, serine, threonine-rich (PEST) region of Mcl–1 (Fig. 1). The five somatic mutations were non-synonymous missense point mutations, as shown in Table II. In addition, four of the mutations were heterozygous (three Mcl–1 mutations and one ASAP3 mutation) and the other was homozygous. Notably, within this small set of genetic alterations, multiple somatic mutations were revealed exclusively in Mcl–1, a member of the B-cell lymphoma (Bcl-2) family. Missense variants in Mcl–1 were present in two of the 20 cases (c.A447G and c.T521C), leading to p.D155G and p.L174S substitutions, respectively. The present study then focussed on Mcl–1 and expanded the investigation into the remaining 43 GBM tissue samples. This revealed another somatic mutation (c.G446C, p.D155H), which was in an identical amino acid position as one of the originally identified somatic mutations in Mcl-1.

Sequence comparisons. As shown in Fig. 1, sequence comparisons demonstrated that the affected amino acids in Mcl-1 are highly conserved among species. Projection of the somatic mutations onto the amino acid sequence of Mcl–1 revealed that all three alterations were located in the PEST region of the enzyme. These three missense point mutations may exhibit functional consequences by affecting the degradation of the protein, since they are all accumulated at highly evolutionarily conserved amino acid residues of the respectively affected subunits (Fig. 1). The PEST region is a well known degradation regulatory element in certain proteins (16). The recurrence of mutations affecting these highly conserved regions involved in the regulation of degradation is suggestive of a gain-of-function effect. Therefore, these mutations are hypothesized to compromise the degradation of the protein, resulting in the intracellular accumulation of Mcl-1.

Mcl-1 post-translational modifications. Mcl-1 is reported to undergo various post-translational modifications and the present study hypothesized that the amino acid alternations identified

<table>
<thead>
<tr>
<th>Gene</th>
<th>Nucleotide</th>
<th>Type</th>
<th>Amino acid change</th>
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<tbody>
<tr>
<td>Mcl-1</td>
<td>GTACGGA&gt;GCGGGTC</td>
<td>Missense</td>
<td>D&gt;D (157aa)</td>
</tr>
<tr>
<td>Mcl-1</td>
<td>GTACGG&gt;CACGGGT</td>
<td>Missense</td>
<td>D&gt;D/H (157aa)</td>
</tr>
<tr>
<td>Mcl-1</td>
<td>ACGAGTT&gt;CTTACC</td>
<td>Missense</td>
<td>L&gt;S (174aa)</td>
</tr>
<tr>
<td>ZMYM3</td>
<td>GGGTCG&gt;CTCTGT</td>
<td>Missense</td>
<td>R&gt;R/P (1363aa)</td>
</tr>
<tr>
<td>SORCS1</td>
<td>GCCGAG&gt;GCGCCT</td>
<td>Missense</td>
<td>G&gt;G/R (460aa)</td>
</tr>
<tr>
<td>ASAP3</td>
<td>TCAATG&gt;CAGGTC</td>
<td>Missense</td>
<td>E&gt;E/Q (504aa)</td>
</tr>
</tbody>
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Mcl-1, myeloid cell leukemia 1; ZMYM3, zinc finger MYM-type protein 3; SORCS1, sortilin-related VPS10 domain containing receptor 1; ASAP3, Arf-GAP with SH3 domain, ANK repeat and PH domain-containing protein 3.
may affect the phosphorylation of the residual amino acids, thereby preventing the degradation of the protein. However, no differences in the phosphorylation of Mcl-1 were identified between the wild-type and the mutants (data not shown). The half-life of the Mcl-1 protein was previously reported to be between 30 and 90 min (16-18). The present study subsequently determined whether the mutations identified affected the half-life of Mcl-1. The wild-type Mcl-1 or the mutant forms were expressed in the GBM cell line and the half-life of Mcl-1 was determined using 35S methionine pulse-chase labeling. Consistent with the previous findings, the wild-type Mcl-1 had a half-life of ~1 h in the U251 cells (Fig. 2). By contrast, the three mutants were more stable compared with the wild-type and exhibited a half-life of ~2 h (Fig. 2).

Cell proliferation. Since Mcl-1 functions as an anti-apoptotic protein and the somatic mutations resulted in the intracellular accumulation of the enzyme, the present study expressed the
mutants and wild-type McI-1 transiently in U251 cells and examined the cell proliferation rate using an MTT assay. As shown in Fig. 3, the ectopic expression levels of the three mutants were significantly higher compared with the wild-type, and the survival rate of the U251 cells transfected with wild-type McI-1 was significantly higher compared with the mutants treated with TRAIL. To investigate the underlying mechanism, the present study examined whether mutant McI-1 had a more marked effect on TRAIL-induced apoptosis in glioma cells using flow cytometry. The McI-1 mutants suppressed TRAIL-induced apoptosis to a greater extent compared with the wild-type in the glioma cell line (Fig. 4).

Discussion

The identification of cancer specific somatic mutations may improve current understanding of the molecular mechanisms underlying tumorigenesis and the progression of malignancy in human cells. This has substantially contributed to the development of highly targeted treatments using monoclonal antibodies or specific inhibitors, which are characterized by a marked improvement in clinical efficacy and an evident reduction in adverse effects compared with conventional chemotherapeutic agents (19,20). Positive results from several clinical trials have led to the approval of similar therapeutic agents for melanoma ( vemurafenib/ BRAF), non-small cell lung cancer (crizotinib/anaplastic lymphoma kinase) and myelodysplasia (ruxolitinib/Janus kinase 2; JNK2) (18), and the improved capability to identify novel targets is likely to assist in expanding the list of targeted therapies. GBM is one of the most life-threatening malignancies and is the most common type of primary brain tumor in adults (21). The standard treatments, including surgery or chemoradiotherapy, are rarely curative, and the majority of tumors recur within a few months. A previous comprehensive genomic study demonstrated that the genetic landscape of GBM is heterogeneous, with 80% of patients affected in one of the three main signaling pathways: p53, phosphatidylinositol-4,5-bisphosphate 3-kinase α and retinoblastoma (6). The present study identified a number of protein-altering somatic mutations by directly sequencing 20 cancer-associated genes (Table II). Notably, within this small set of genetic alterations, multiple somatic mutations were identified exclusively in McI-1. Missense variants in McI-1 were present in two of the 20 cases (D155G and L174S). Investigating McI-1 in the remaining 43 GBM cases, revealed that three alterations were located in the PEST region of the enzyme. The recurrence of mutations affecting these highly conserved regions, which are involved in the regulation of degradation, is suggestive of a gain-of-function effect. To investigate the effect of these somatic mutations on the function of McI-1, the coding sequences of wild-type McI-1 and the mutants corresponding to the amino acid alterations were cloned into pcDNA4. These constructs were subsequently expressed in the GBM cell line and the half-life of McI-1 was determined by 35S methionine pulse-chase labeling. Wild-type McI-1 exhibited a half-life of ~1 h in the U251 cells (Fig. 2). By contrast, the mutants were more stable, with a half-life of ~2 h (Fig. 2).

McI-1 is readily induced by various survival regulators, including epidermal growth factor, vascular endothelial growth factor, granulocyte-macrophage colony-stimulating factor, mitogen activated protein kinase and JAK/STAT signal transducer and activator of transcription signaling cascades, and is also rapidly degraded by certain apoptosis-inducing signals (32,33). The rapid induction and degradation of McI-1 suggested that McI-1 functions as sensor and reactor of environmental stimuli to maintain a balance between cell survival and death, suggesting that McI-1 is an essential regulator of proliferation or differentiation of human cells (34,35). Inhibition or elimination of McI-1 in response to cytotoxic signals is considered critical in cell death in a number of normal and malignant cells (34,36). In addition, increased expression of McI-1 may not only promote short-term survival in a wide range of cells, but may also cause long-term immortalization and the malignant transformation of certain human cells (37-39). Previous studies have suggested that overexpression of McI-1 may be involved in the mechanism underlying chemoresistance in a number of human malignancies, including breast cancer, leukemia, melanoma, pancreatic cancer and hepatocellular carcinoma (33,40). The present study examined the effect of the identified mutants on the growth of glioma cells and TRAIL-induced apoptosis in U271 cells, and found that the survival rate of the cells transfected with wild-type McI-1 was significantly lower compared with the McI-1 mutants. Additionally, the McI-1 mutants suppressed TRAIL-induced apoptosis to a greater extent compared with the wild-type in the glioma cell lines (Figs. 3 and 4).

In conclusion, despite the small sample size, the present study identified a panel of somatic mutations in 20 cancer-associated genes, and found that several mutations may be pathogenic with potential detrimental impacts on gliomagenesis, within which multiple somatic mutations were identified in McI-1. The preliminary functional investigations suggested that the mutations increased the stability of McI-1 and contributed to the pathogenesis of GBM. Future studies using a larger number of patients from different ethnic backgrounds are required to confirm the findings of the present study.


McL-1 is phosphorylated in the PEST region and stabilized upon ERK activation in viable cells and at additional sites with cytotoxic okadaic acid or taxol. Oncogene 23: 5301-5315, 2004.


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