Role of metallothionein 1E in the migration and invasion of human glioma cell lines

HYANG-HWA RYU23, SHIN JUNG12, TAE-YOUNG JUNG12, KYUNG-SUB MOON1, IN-YOUNG KIM1, YOUNG-II JEONG2, SHU-GUANG JIN12, JIAN PEI12, MIN WEN12 and WOO-YEOL JANG1

Departments of 1Neurosurgery, 2Brain Tumor Research Laboratory, Chonnam National University Research Institute of Medical Sciences, Chonnam National University Hwasun Hospital and Medical School, Gwangju; 3Department of Clinical Laboratory Science, Seonam University, Namwon, Republic of Korea

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Abstract. Metallothionein 1E (MT1E) has been found to be highly expressed in motile cell lines. We investigated whether MT1E actually modulates the migration and invasion of human glioma cell lines and the types of factors that have an effect on MT1E. RNA differential display was performed using GeneFishing™ technology in the human glioma cell lines U343MG-A, U87MG and U87MG-10; the results were validated by RT-PCR and northern blot analysis, in order to detect possible genetic changes as the determining factors for migration ability in malignant glioma. MT1E was identified in U87MG, a highly motile cell line. The migration and invasion abilities of human glioma cell lines, and MT1E transfectants were investigated using simple scratch testing and Matrigel invasion assays. Morphological and cytoskeletal (actin, vimentin) changes were documented by light and confocal microscopy. The expression of MT1E in four glioma cell lines was assessed by RT-PCR and western blotting. In addition, the effects of MT1E on the activity of the NF-κB p50/p65 transcription factor, MMP-2 and -9 were examined by western blotting and zymography. The endogenous MT1E expression in the human glioma cell lines was statistically correlated with their migratory abilities and invasion. The U87-MT-AS cells became more round and had decreased stress fibers, compared with the U87MG cells. Endogenous MT1E expression in the four human glioma cell lines was directly correlated with migration. Two antisense MT1E-transfected cell lines showed decreased NF-κB p50 translocation into the nucleus, which led to decreased activity of MMP-9 in conditioned media. It may be postulated that MT1E can enhance the migration and invasion of human glioma cells by inducing MMP-9 inactivation via the upregulation of NF-κB p50.

Introduction

The failure of surgical treatment in patients with glioma is mainly due to the invasion of tumor cells into the normal brain beyond the resection areas. These invasive remaining cells are inoperable, resistant to radio- and chemotherapy, and eventually lead to tumor regrowth. Finally, due to the invasion ability of malignant glioma, patients generally die within 1 year after the initial diagnosis (1,2).

To infiltrate into adjacent tissues, certain tumor cells should be able to recognize the barrier matrix, to breach the matrix and to grow into the ectopic locale. Other tumor cells also need to control signaling pathways that would contribute to tumor invasion. It is difficult to directly investigate the specific phenotype that is associated with invasion ability in vivo. The dysregulation of the induced motility of tumor cells has been postulated to be one of the invasion processes that functions in the transmigration of the barrier matrix. Thus, there is a good correlation between increased invasiveness in vivo and increased motility in vitro (1,3,4). The identification of genes related to increased motility is critical for understanding the molecular basis of biological behavior in invasive gliomas.

In a previous study, the authors sought to identify genes that could serve as determining factors for the mobility of malignant glioma cells using DD-PCR (5,6). The results showed that metallothionein 1E was highly expressed in motile cell lines.

Metallothionein 1E (MT1E) is the name for a family of low-molecular weight intracellular metalloproteins with a high affinity for heavy metal ions, such as zinc, cadmium, copper, mercury, and platinum. A striking feature of these metalloproteins is that they are involved in heavy metal detoxification, chemoresistance to anti-cancer drugs and free radical scavenging for cell protection (7-11). Furthermore, they can modulate the activities of zinc-dependent regulatory proteins including enzymes and zinc-finger transcription factors, by the removal and transfer of zinc (12,13). It is also well known that MT participates in fundamental cellular processes, such as cell proliferation and apoptosis (14,15). However, even though many reports have been issued concerning MT, its functions...
associated with the migration and invasion have not reported in glioma.

Nuclear factor κB (NF-κB) is a transcription factor that regulates an exceptionally large number of genes, particularly those involved in immune and inflammatory responses. NF-κB is a dimer most commonly composed of two protein subunits, p50 and p65, and normally remains in an inactive form in the cytoplasm bound to an inhibitor subunit, IκB. Various stimuli activate upstream IκB kinases, which in turn phosphorylate and degrade IκB. The released NF-κB then enters the nucleus and activates the transcription of many different target genes. NF-κB also plays a pivotal role in a diverse array of cellular activities associated with the regulation of cell death, growth, and development (16). NF-κB has far reaching importance in the regulation of cell death as the overexpression of NF-κB renders cancer cells resistant to chemotherapeutic agents (17-19).

MT1E may act as a potential intracellular modulator of NF-κB activation by regulating the cellular level and activity of NF-κB (20,21). Despite the obvious importance of understanding the MT1E-NF-κB interaction, little is known about the tentative molecules that coordinate this interaction.

The relationship between tumor cells and host factors have an effect on invasion, thus a greater understanding of these factors could be important in the anti-invasive research. Primary tumor cell invasion of surrounding tissue is the first stage of a metastasis cascade, and many factors, such as matrix metalloproteinases (MMPs), are associated with this stage (22). MMPs belong to a family of zinc-dependent endopeptidases that degrade the extracellular matrix. Especially, MMPs have well known effects on tumor proliferation and metastasis. Therefore, we investigated whether MT actually modulates migration and invasion in human glioma cells by modulating activity of the NF-κB, MMP-2 and -9.

Materials and methods

Cell lines and cell culture. Human glioma cell lines, U87MG, U251MG, U373MG and U343MG-A were obtained from the Korean Cell Line Bank, Seoul, Korea and from the Brain Tumor Research Center, University of California, San Francisco, CA, USA, respectively. U87MG-10' was obtained by repetition scratch to tenth U87MG. All cell lines were routinely grown in Dulbecco’s modified Eagle’s medium (Gibco-BRL, Gaithersburg, MD, USA) supplemented with 10% fetal bovine serum (Gibco-BRL, Gaithersburg, MD, USA) at 37°C in a humidified 95% air/5% CO₂ atmosphere. Samples of astrocytic tumors were obtained from the patients with the surgery performed at Chonnam National University Hospital. The specimens were snap-frozen in liquid nitrogen and stored at -70°C until use.

Migration assay. In order to stop cell proliferation, the media used to culture cells were replaced with medium containing 5 mM hydroxyurea. After 24 h of hydroxyurea treatment, the cultures were scraped with a single-edged razor blade. Cells were washed twice with PBS and placed in medium containing hydroxyurea. After 48 h of incubation, the cells were washed twice with PBS, fixed with absolute alcohol, and stained with 0.1% toluidine blue. Six microscopic fields were evaluated for each wound injury. The number of cells migrating across the wound edge and the maximum distance migrated were determined in each field and averaged for each injury. These experiments were repeated three times.

RNA isolation from glioma cell lines. Total RNA was isolated from the malignant glioma cell lines using TRIzol (Life Technologies, Gaithersburg, MD) according to the manufacturer’s instructions. After isolating the RNA, the RNA pellets were frozen and stored at -80°C until use.

Differentially display-polymerase chain reaction (DD-PCR). DD-PCR was performed using the Genefishing™ kit (Seegene Incorp., Seoul, Korea) according to the manufacturer’s instructions. For first strand synthesis, 3 μg of the purified total RNA was incubated with 10 μM dT-ACP for 3 min at 80°C, followed by the addition of a buffer containing 4 μl of 5X RT buffer (Promega, Madison, WI, USA), 1 μl of dNTP (2 mM each), 2.5 μl of 25 mM MgCl₂, 2 μl of RNase Inhibitor (40 U/μl Promega), and 1 μl of reverse transcriptase (200 U/μl Promega). The reaction volume was 20 μl, and the reaction was allowed to proceed for 90 min at 42°C, and then for 2 min at 90°C. The PCR protocol for second-strand synthesis was one cycle at 94°C for 3 min, followed by 50°C for 3 min, and 72°C for 1 min. After second-strand DNA synthesis was completed, the PCR products were subjected to second-stage PCR amplification, which included 40 cycles of 94°C for 40 sec, 65°C for 40 sec, 72°C for 40 sec, followed by a 5 min final extension step at 72°C. The PCR products were separated on a 1.2% agarose gel and stained with ethidium bromide. The differentially expressed bands were extracted from the gel using a QIAquick Gel extraction kit (Qiagen, Carlsbad, CA, USA), directly cloned into the pGEMR-T Easy vector (Promega) without reamplification of the recovered bands, and sequenced.

Reverse transcription polymerase chain reaction (RT-PCR). Total RNA was isolated from three cultured glioma cell lines using TRIzol reagent. A total of 1 μg of RNA was reverse transcribed to synthesize the cDNA. Reverse transcription (RT) was followed by polymerase chain reaction (PCR). For the first strand synthesis, 1 μg of the purified total RNA was incubated with oligo (dT) (0.5 μg/μl Promega) for 5 min at 70°C, followed by the addition of buffer containing 4 μl of 5X Reaction buffer (Promega), 1 μl of dNTP (10 mM each), 3.5 μl of 25 mM MgCl₂, 2 μl of RNase Inhibitor (40 U/μl Promega), and 1 μl of reverse transcriptase (200 U/μl Promega). The reaction volume was 20 μl, and the reaction was carried out for 90 min at 42°C, and then for 2 min at 90°C. β-actin was used as an internal control. Reactions with varying numbers of polymerase chain reaction (PCR) cycles were run for each transcript.

Northern blot analysis. Northern blot analysis was performed under conventional conditions. Total RNA (20 μg) was separated by electrophoresis on 1.2% agarose formaldehyde gels, then transferred to a nylon membrane overnight and cross linked with UV irradiation. The filters were prehybridized at 65°C for 3 h and then hybridized to a 32P-labeled probe. The probes were prepared from the clones described above by digestion with restriction enzymes, and this was followed by gel electrophoresis. The filters were washed in SSC and then 0.1% SDS for 15 min at 65°C. They were then exposed to X-ray film at -70°C.
Transfection

Preparation of metallothionein IE construct. The complete coding region of the human MT cDNA was amplified from U87MG cDNA by PCR with synthetic primers. The PCR primers were designed as follows. MT-sense: 5′-CGGATCCATGGACCCCAACTGCTCTT-3′, 5′-GCTCTAGAAGCTCAGGCACACAGCCTGAC-3′. MT-anti sense: 5′-GCTCTAGAGCATTGACCCCAACTGCTCTT-3′, 5′-CGGGATTCCTGACACAGCCTGAC-3′. The amplified cDNA was sequenced and directly subcloned into the pcDNA3.1+(+) vector (Invitrogen, San Diego, CA, USA) between the BamHI and XhoI sites, which is contained with in the CMV promoter and the neomycin resistance gene. The resulting vectors were designated as pcDNA3.1-MT-S, pcDNA3.1-MT-AS.

Transfection

U343MG-A and U87MG were maintained under exponential growth conditions in DMEM supplemented with 10% fetal bovine serum in the absence of antibiotics. The optimal cell density for transfection is normally between 50-80% confluency for adherent cells. pcDNA3.1-MT-S/AS were respectively transfected into human malignant glioma cell lines U343MG-A and U87MG using the GeneJuicer™ transfection reagent (Novagen, Madison, WI, USA). Cells in serum-free DMEM were mixed with 15 µg of plasmid DNA and 45 µl of GeneJuicer-serum-free media according to the manufacturer’s protocol. After incubation at 37°C (5% CO₂) for 5 h, the transfection mixture was replaced with DMEM supplemented with 10% FBS. After 24 h incubation, the medium was replaced with DMEM containing 10% FBS and 500 µg/ml G418. U343MG-A transfected with sense MT1E cDNA plasmid (pcDNA3.1-MT-S) and U87MG transfected with antisense MT1E cDNA plasmid (pcDNA3.1-MT-AS). Each transfectant was designated as U343-MT-S and U87-MT-AS.

Separation of cytoplasm and nuclear extract. Cytoplasmic extraction buffer (CEB) consisted of 10 mM HEPES (pH 7.9), 40 mM KCl, 2 mM MgCl₂, 10% glycerol and protease inhibitors. After the cells were scraped into CEB, they were transferred into a micro-centrifuge tube, lysed by vortexing a few times and chilled on ice for 15 min. Nuclei were pelleted by centrifugation at 5,000 rpm for 5 min, and the pellets were washed twice with PBS. The supernatant was removed, and the nuclei were lysed in nuclear extraction buffer consisting of 10 mM HEPES (pH 7.9), 500 mM NaCl, 1% Triton X-100, 10% glycerol and protease inhibitors. After centrifugation at 12,000 rpm for 5 min, the supernatants containing the nuclear proteins were stored at 20°C until use.

Preparation of total protein and conditioned media. For the preparation of total protein, cells were lysed in a protein extraction buffer [50 mM Tris (pH 8.0), 5 mM EDTA, 150 mM sodium chloride, 0.5% deoxycholic acid, 0.1% SDS, 1% NP-40, 1 mM PMSF, and 1 mg/ml protease inhibitor cocktail].

For the preparation of conditioned media, cells were grown in 60-mm plates until they were subconfluent, and were then washed three times with ice-cold CMF-PBS. 1 ml of serum-free medium was added to each plate, and the cells were incubated at 37°C for 48 h. The conditioned media were clarified by centrifugation. The concentrations of all kinds of protein samples were determined using the Bio-Rad protein assay kit (Bio-Rad, Hercules, CA, USA).

Western blotting. A total of 20 µg of whole cell lysates were separated by 15% SDS-PAGE and transferred to a polyvinylidene difluoride membrane (Pall corporation, USA). The membrane was then incubated for 2 h at room temperature in TBS-T solution [10 mM Tris-Cl (pH 8.0), 150 mM NaCl, and 0.05% Tween 20] supplemented with 5% non-fat dry milk and probed overnight at 4°C with anti-metallothionein (Santa Cruz Biotechnologies, Santa Cruz, CA, USA), anti-p50, p65 (Cell Signaling Technology, Beverly, MA), anti-MMP2 and MMP9 (BD Pharmingen, San Diego, CA, USA). The bound antibodies were visualized with the appropriate horseradish peroxidase-conjugated secondary antibody (Jackson Immunoresearch Laboratory, WestGrove, PA, USA) conjugated with horseradish peroxidase using enhanced chemiluminescence reagents (ECL, Amersham Biosciences, Bucks, UK).

Cell morphological characteristics. The transfectants and the parental cells were fixed with methanol at 4°C for 10 min. Cells were washed twice with PBS, fixed with cold methanol and stained with 0.1% toluidine blue. These were examined via light microscopy (Nikon, Garden City, NY) and digitally photographed to evaluate and record the morphology of the cell population.

Doubling time of stable transfectants. The cells were seeded at 3x10⁴ cells in 60 mm culture dish. After serum starvation for 24 h, the cells were counted each day. The cells were treated with trypsin-EDTA and the number of viable cells was counted with a hemocytometer. The doubling-time was calculated from the cell growth curve over 5 days. Equation for doubling time = (t₀-t)log₂/logN₀/N (t, final time; unit, hour; t₀, initial times; N, final cell number; N₀, initial cell number).

Immunocytochemistry for cytoskeletal proteins. The cells were cultured on glass coverslips in 35 mm dishes until reaching subconfluency, washed with phosphate buffer saline and fixed with 4% p-formaldehyde for 10 min. After washing in immuno/DNA buffer (Research Genetics, Huntsville, AL, USA), the cells were treated with trypsin-EDTA and the number of viable cells was counted with a hemocytometer. The doubling-time was calculated from the cell growth curve over 5 days. Equation for doubling time = (t₀-t)log₂/logN₀/N (t, final time; unit, hour; t₀, initial times; N, final cell number; N₀, initial cell number).

Invasion assay. Matrigel (reconstituted basement membrane; 25 µg) was dried on a polycarbonate filter (polystyrene-pyrrolidone-free; Nucleopore; Whatmann, Maidstone, UK). Cells...
were harvested by brief exposure to 1 mM/l EDTA, washed with DMEM containing 0.1% bovine serum albumin, and added to the Boyden chamber (2x10^5 cells). Cells were incubated for 24 h at 37˚C in a humidified atmosphere of 95% air and 5% CO₂. Cells that transversed the Matrigel layer and attached to the filter were stained with the Diff Quik kit (Dade Diagnostics, Aguada, PA, USA) and counted in five randomized fields. The results are expressed as the mean ± SE of three independent experiments.

Data analysis. The comparison of the nucleotide sequence homology of the isolated cDNAs with the registered sequence in Genebank was carried out using the BLAST algorithm. We measured the statistical significance of the cell distance and the cell number using the Mann-Whitney U-test, and the doubling time by repeated measures ANOVA. *P*<0.05 was considered to be significant. Statistical analysis was performed using the SPSS software program (version 12.0 for Windows; SPSS Inc., Chicago, IL).

Results

Comparison of the migration abilities of three glioma cell lines. The motility of the cell lines was compared using a simple scratch technique. In order to eliminate any confounding effects of an experimental agent on cell proliferation, the cell culture media were replaced with medium containing hydroxyurea. Treatment with hydroxyurea resulted in the complete inhibition of cell proliferation. As shown in Fig. 1, the U87MG cells were more motile than the U343MG-A cells, and the U87MG-10' cells were the fastest. The difference among the three groups was statistically significant for the mean cell number (p<0.021).

Identification of differentially expressed genes. To obtain a profile of the genes related to invasion, the author estimated the motility of the three cells lines that showed different motilities.
MT1E showed a higher expression level in the motile cell lines (Fig. 2A).

Validation of DD-PCR data by RT-PCR and northern blot analysis. The author verified the significant differences in gene expression pattern by RT-PCR and northern blot analysis for the candidate gene. As shown in Fig. 2B and C, there were significant differences in MT expression between the low and high motility cell lines, and the expression patterns matched those of both RT-PCR and northern blot analysis.

Selection of stable transfectants of sense and antisense MT constructs. To assess the effect of MT on proliferation, migration and invasion in malignant glioma cells, the pcDNA 3.1(+) vector containing full length sense and antisense MT1E cDNA was transfected into malignant glioma cell lines under the control of the CMV promoter, and neomycin-resistance transfected clones were obtained. U343MG-A cells expressing low levels of MT1E were inserted into the sense MT1E cDNA construct. In addition, U87MG cells expressing high levels of MT1E were inserted into an antisense MT1E cDNA construct. Western blotting was used to detect the clone that showed the best characterization after transfection with sense and antisense constructs. The U343-MT-S cell line showed a strong band when compared with the parental cell line (Fig. 3A). On the other hand, the U87-MT-AS cell line showed a very faint band when compared with parental cell line (Fig. 3B).

The effect of MT on transfectants of sense and antisense MT constructs. In order to observe changes in cell motility alone. As shown in Fig. 4A, MT1E sense/antisense-transfected cell lines showed positively altered their motilities. U343MG-A is a less motile cell line. Following sense MT1E transfection, it becomes about 37% faster than parental cell lines. The maximum distance migrated and the mean cell number was 261 µm and 78 cells in the U343MG-A cell line and 358 mm and 148 cells in the U343-MT-S cell line, respectively. The difference between the two groups was statistically significant for the mean cell number (p<0.001) and maximal distance (p<0.001). Likewise, U87MG is a high motility cell line, and it became less motile after transfection with antisense MT. The maximum distance migrated and the mean number of total cells migrated were 844 µm and 495 cells in the U87MG cell line and 659 µm and 270 cells in the M87-MT-AS cell line, respectively. The difference between the two groups was also statistically significant for the mean cell number (p<0.001) and maximal distance (p<0.001). In addition, MT1E on the invasion in the sense and antisense transfectants was markedly enhanced in the Matrigel assay as compared with the parental cell line (p<0.001, Fig. 4B).

Morphological and cytoskeletal changes after MT transfection. Morphological changes were characterized by light microscopy. U343MG-A cells generally formed colonies and were round in shape. On the other hand, U343-MT-S cell line grew apart from each other, and their shape became more spindle-like (Fig. 5A and B). In addition, U87-MT-AS cells became flat and enlarged in comparison to U87MG cells (Fig. 5C and D). Immunofluorescence staining of actin and vimentin were performed in order to determine whether the differences in cytoskeletal alterations were associated with tumor cell motility and invasion. The expression patterns of actin and vimentin were similar. The U343-MT-S and U87MG cell lines with high motility formed many stress fibers and showed extensive lamellipodia and strong positive staining in comparison with the parental cell lines. The proteins were distributed on one side...
around the nucleus and cytoplasm (Fig. 6F and I). On the other hand, the U343MG-A and U87-MT-AS cell lines, which were less motile, showed fewer stress fibers and shorter lamellipodia. The cytoskeletal proteins were mainly concentrated around the nucleus and became entangled (Fig. 6C and L).

**Doubling time test.** The growth rate was compared of each cell line with those of transfected clones. As shown in Fig. 7, the doubling time was 35 and 31 h in the U343MG-A and U343-MT-S cell lines, respectively. The doubling time for the U87MG and U87-MT-AS cell lines was 32 and 39 h, respectively. U343-MT-S cell line was reduced by about 4 h and that of the U87-MT-AS cell line was only about 7 h, compared with parental cell lines.

**Expression of endogenous MT in malignant glioma cell lines.** RT-PCR was used to confirm the expression of MT1E in glioma cell lines with different motilities. The U87MG, U251MG, U373MG and U343MG-A cell lines showed high motility sequences (Fig. 8A and B). The levels of MT1E mRNA were proportional to the motility of the cell lines (Fig. 8C).

**Effects of MT on NF-κB translocation following the activity of MMP-2 and -9.** U87MG cells were stably transfected with antisense MT1E cDNA, and the best two clones were selected (U87-MT-AS). In order to examine the relationship between MT1E and NF-κB activity, the p50 and p65 subunits were detected in the nuclear and cytoplasm fraction by western blot analysis. As shown in Fig. 9A-b, U87-MT-AS showed decreased translocation of the p50 subunit of NF-κB from the cytoplasm to the nucleus when compared with the parental cells. However, the translocation of NF-κB p65 was not particularly changed in comparison with the parental cells. In addition, total IκB was increased in U87-MT-AS (Fig. 9A-a). On the contrary, total IκB was only slightly increased in U343-MT-S. Thus, we checked phospho-IκB expression level in U343-MT-S. As shown in Fig. 9B, the expression of phospho-IκB was increased with total IκB. The expression of MMP-2 in U87MG and U87-MT-AS were detected by western blotting and zymography. But the expression of MMP-9 was only confirmed by western blotting because it could not be detected by zymography. There was no difference in MMP-2 expression between U87MG and U87-MT-AS. However, the active and latent forms of MMP-9 were remarkably decreased in the conditioned media of U87-MT-AS (Fig. 9A-a), otherwise U343-MT-S had increased active forms of MMP-9 in conditioned media.

**Discussion**

In a previous *in vitro* study, we identified motility-related genes that would be representative of *in vivo* invasiveness in glioma cell lines with different motilities using the Genefishing technology (6). Among the identified genes, MT1E was overexpressed in highly motile malignant glioma cell lines, indicating that this gene may be a positive regulator of tumor cell motility. In this study, the author investigated whether MT1E is actually associated with migration and invasion in human glioma cell
lines. Firstly, two glioma cell lines were selected with similar origin, but showed a marked difference of their motility (Fig. 4). Then a sense MT1E cDNA construct was transferred into the glioma cell line with less motility, and an antisense MT1E cDNA construct was transferred into glioma cell line with high motility. The effect of MT1E on the migratory and invasive abilities of human glioma cell lines was evaluated by an in vitro assay that involved a simple scratch technique and the Boyden chamber system. The results showed that the proliferation rate in the U343-MT-S cell line, an MT1E-overexpressing cell line, was two times faster, whereas the opposite effect was observed in the U87-MT-AS cell line, which expressed lower levels of MT1E. In addition, some malignant glioma cell lines with different motilities expressed MT in proportion to their mobility. Generally, cell movement is accompanied by various changes, such as morphology, cytoskeleton, and proliferation rate. To determine whether the differences associated with cytoskeletal alterations were also associated with tumor cell motility, immunofluorescence staining for actin and vimentin was performed and the morphological changes in the shape and size of the cells following transfection were examined. The U343-MT-S cells grew apart from each other and showed well-defined stress fibers. On the other hand, when this cell line was transfected with antisense MT1E, the cells became flatter and larger, and the cytoskeletal proteins became entangled. In the process of several experiments, the antisense transfectants was more sensitive to trypsin than the parental cell line (data not shown), while the sense transfectants were resistance to it. This may indicate that these cells are ready to migrate.

MT is related to tumor grade, malignancy and prognosis in certain tumors. MT is correlated with the histological grade and proliferative potential of astrocytic tumors (23,24). In addition, MT is overexpressed in invasive and in situ breast cancer cells, and it may be a biomarker of tumor differentiation and aggressiveness (25,26). In order to verify whether MT1E affects the proliferation of glioma cell lines, doubling time test in U343-MT-S and U87-MT-AS, was performed compared with each parental cell line. Actually, U343-MT-S increased cell proliferation rate, while U87-MT-AS was the opposite. On the basis of the above results, MT-1E may be directly or indirectly involved in cell motility and invasion.

Structural studies of MT have shown that this unusual protein with 61 amino acids (mammalian MT) can bind with both essential metals (zinc and copper) and toxic metals (cadmium and mercury) in two distinct cluster structures in the molecule (27). Due to this characteristic, MTs act as controllers of zinc and copper, by participating in cell proliferation processes (28). MT could donate zinc/copper to various metallo-enzymes and transcription factors (29,30). The protective role of MT in oxidative stress and metal toxicity suggests that MT may also have a functional role in tumor cell survival and growth. According to some reports, MT could facilitate tumor cell growth by two potential mechanisms. One possible mechanism is that MT may act as a zinc donor to various transcription factors, including tumor suppressor gene products, such as p53. In vitro studies have shown that thionein can modulate the transcriptional activation of Sp1, a zinc finger transcription factor. Another possible mechanism is that MT may protect the cells from radiation and
chemotherapeutic agents by virtue of its free radical scavenging property (31-33). We placed major emphasis on first hypothesis, and then desired to look for any factors which have Zn associated site in their structure and associated in tumor cell invasion. It may be related with MT1E in the change of the motility of malignant glioma cell lines.

MMPs are a part of a family of Zn-dependent enzymes and have a well known activity in tumor invasion and metastasis (39). The change of the expression level of MMP-2 and -9 was examined in both total cell lysates and conditioned media in U87MG and U87-MT-AS. The activity of MMP-9 was decreased in U87-MT-AS, while the activity of MMP-2 was similar, in comparison to the parental cell line. In 1996, Haga et al reported that metallothionein increased the activity of MMP, which may change in conformation by chelation with metallothionein (40).

NF-κB is also a zinc-dependent transcription factor, plays a pivotal role in a diverse array of cellular activities and gene activations (30). There are strong pathophysiological functional similarities between MT and NF-κB. Both are anti-apoptotic entities, especially in cancer cells and both have regulatory roles in inflammation (34,35). Moreover, many stimulators of their expression overlap, including tumor necrosis factor-α, lipopolysaccharide, and interleukin-1 (36). Thus, it is plausible that there is a close molecular relationship between the two molecules. A previous study showed that MT regulates the cellular level and activity of NF-κB as a potential intracellular modulators of NF-κB activation. In addition, MT overexpression was found to up-regulate DNA binding of the NF-κB (21,31,37), and it plays an important role in MMP-9 gene expression (38).

In this study, when MT1E expression was decreased in glioma cell line with high motility, the levels of NF-κB p50 in the nuclear fraction significantly decreased. Most reports on the association between MT and NF-κB have suggested that it involves the p65 subunit of NF-κB. However, the present study showed that MT affected the p50 subunit of NF-κB rather than the p65 subunit. A 1998 report suggested that MT specifically interacts with the p50 subunit of the p50/RelA heterodimer and that MT may be required for the stabilization of its DNA complex formation, thus allowing for the potent transactivation of target genes (39). According to the last report, MT2A promotes breast cancer cell invasion by upregulating MMP-9 via AP-1 and NF-κB (41).

On the basis of our results and reports, MT1E can modulate the motility and invasion in a human glioma cell line. It may involve two mechanisms. MT1E may induce the change of morphology via the modification of cytoskeletal proteins in human glioma cell lines, or MT1E may modulate the activity of MMP-9 by NF-κB in malignant glioma cell lines. This series of events enhances migration and invasion in malignant glioma cell lines. More experiments should be performed to clarify these mechanisms.

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