SUPPRESSION OF MANGANESE SUPEROXIDE DISMUTASE ACTIVITY IN ROTENONE-TREATED HUMAN GliOBLASTOMA T98G CELLS REDUCES CELL VIABILITY

SEPTELIA INAWATI WANANDI1*, NOVI SILVIA HARDIANY1, NURJATI CHAIRANI SIREGAR2, MOHAMAD SADIKIN1

1Department of Biochemistry and Molecular Biology, Faculty of Medicine, Universitas Indonesia, Jakarta, Indonesia. 2Department of Anatomic Pathology, Faculty of Medicine, Universitas Indonesia, Jakarta, Indonesia. Email: septelia@gmail.com/septelia.inawati@ui.ac.id

ABSTRACT

Objective: Glioma is the most common human primary brain tumor which is highly resistant to oxidative stress-based anticancer. The aim of this study was to analyze the effect of rotenone-induced reactive oxygen species (ROS) on the modulation of manganese superoxide dismutase (MnSOD) expression and cell viability in human glioblastoma (GBM) T98G cells.

Methods: In this in vitro experimental study, T98G cells were treated with high-dose rotenone (0.5, 5, and 50 µM, respectively). Following rotenone treatment and intracellular ROS, both peroxide and superoxide radicals were determined. Moreover, we analyzed MnSOD mRNA expression, protein, and specific activity, as well as cell survival including viability, proliferation, apoptosis, and mitochondrial structure.

Results: High-dose rotenone treatment of T98G cells significantly increased intracellular ROS and MnSOD mRNA, but its protein and specific activity definitely decreased. The treatment also led to a reduction of cell viability, enhancement of apoptosis, and disruption of mitochondrial integrity.

Conclusion: Overproduction of ROS in rotenone-treated human GBM T98G cells could suppress the MnSOD protein level and activity even though mRNA synthesis has been increased. This modulation led to reduced survival of T98G cells through induction of cell death rather than inhibition of cell proliferation.

Keywords: Cell viability, Glioblastoma multiforme, Manganese superoxide dismutase, Rotenone.

INTRODUCTION

Glioma is the most common human primary brain tumor which arises from glial cells [1]. Nowadays, the use of conventional treatments such as chemotherapy and radiation does not significantly enhance the life expectancy of glioma patients, particularly those with high-grade malignant glioblastoma (WHO Grade IV) patients, which is highly resistant to therapy [2]. Combinatorial treatment strategy has been reported to have improved the chemotherapeutic delivery to tumor cells in the brain [3,4]. Nevertheless, accumulating evidence suggests that dysregulation of cell cycle and apoptosis could lead to radiotherapy resistance [5]. A plausible mechanism for this resistance might involve high antioxidant status in tumor cells which could affect cell survival in response to radiation-induced oxidative stress [6].

Manganese superoxide dismutase (MnSOD) is a major cellular antioxidant located in the mitochondrial matrix. This enzyme catalyzes the reaction of superoxide anion into hydrogen peroxide which will be later eliminated by catalase or peroxidase. The previous studies have reported that the upregulation of MnSOD expression could inhibit the phenotype of various cancer cells, suggesting that MnSOD is a tumor suppressor [7-10]. However, this assumption is still controversial, since it has been demonstrated that MnSOD was overexpressed in several human cancers including GBM [11]. Furthermore, MnSOD has been reported to have an important role on tumor cell growth and proliferation in ovarian cancer through regulating no superoxide level [12]. Our recent study has confirmed that MnSOD mRNA expression and specific activity in human glioma cells isolated from clinical specimens were higher than those in normal brain cells. In addition, the oxidative stress biomarkers, i.e., malondialdehyde, carbonyl compounds and 8-OHdG, were notably enhanced [8]. Interestingly, when these cells were categorized based on tumor grade, we found that the high-grade malignant glioma cells expressed MnSOD mRNA at higher levels compared to the low-grade but had lower specific activity.

The differential between MnSOD mRNA level and specific activity in high-grade glioma prompted a line of inquiry to investigate whether this was associated with reactive oxygen species (ROS) levels in these cells. Therefore, the aim of this study was to analyze the effect of rotenone-induced ROS in human GBM on the modulation of MnSOD expression and its association with cell viability. High-dose rotenone was applied to human GBM T98G cells to induce overproduction of intracellular ROS, which is an expected consequence of radiotherapy. Rotenone is an inhibitor of mitochondrial complex I electron transport chain, which increases the mitochondrial ROS generation, particularly superoxide radicals, and leads to cytotoxicity [13-15]. It has also been reported that this toxin could be used as an anticancer agent [15]. Finally, this study is also projected to elaborate on the role of MnSOD in high malignant GBM patients who are resistant to radiotherapy. The outcome of this study has the potential for improving the management of GBM.

MATERIAL AND METHODS

Cell culture

The human GBM cell line T98G (kindly provided by Prof. Alexander Brehm from Institut fuer Molekularbiologie und Tumorforschung Philippus Universitat Marburg, Germany; ATCC No. CRL-1690™) was maintained in high glucose DMEM containing 10% of heat-inactivated fetal bovine serum, 3.7 g/L of sodium bicarbonate, 1% Streptomycin - Penicillin, and 1% amphotericin B at 37°C in a humidified atmosphere of 95% air and 5% CO2. The cells were subcultured with 0.25% trypsin and 1% ethylenediaminetetraacetic acid (EDTA) whenever the culture reached confluence. A subcultivation
ratio was 1.2–1.5. Cells were utilized for analysis within 15 passages since it has been demonstrated that antioxidant enzyme levels in tumor cells subcultured up to 50 passages did not change [8].

**Rotenone treatment**

Rotenone powder (Sigma Aldrich, USA) was first dissolved in dimethyl sulfoxide (DMSO) to obtain 10 mM stock solution. One day before rotenone treatment, 5 × 10^4 T98G cells were plated triplicate in a 24-well plate and grown in the same medium without free serum. Cells were treated with various concentrations of rotenone and incubated for 6 h, as described previously [12]. Negative controls were T98G cells without rotenone or DMSO treatment, as well as T98G cells treated with the same volume of DMSO used to dissolve rotenone. Rothenone treatment was repeated 3 times at different time points. After the treatment, cells were washed with phosphate-buffered saline (PBS), harvested and then centrifuged at 1000 rpm for 5 min. Finally, cells were counted and prepared either for the cell viability or proliferation analysis, as well as for the isolation of total RNA and protein, followed by the analysis of MnSOD mRNA expression, protein level, and specific activity, respectively.

**Determination of intracellular ROS level**

Intracellular ROS was measured at the level of hydrogen peroxide and superoxide anion. For peroxide level measurement, dichlorodihydrofluorescein diacetate (DCFH-DA) assay (Molecular Probes, USA) was performed based on the intracellular peroxide-dependent oxidation of 2',7'-DCFH-DA to form the fluorescent compound 2',7'-dichlorofluorescein DCF. As described previously [16], Dihydroethidium (DHE) (Invitrogen) assay could detect the presence of superoxide anion which enzymatically converts hydroethidine to ethidium. T98G cell pellets containing 2×10^4 cells were resuspended in PBS containing 20 μM DCFH-DA or 20 μM DHE and incubated for 30 min at 37°C [17,18]. Cells were then rinsed with PBS (2 times) and centrifuged 1000 rpm for 5 min. Cell pellets were resuspended in 3 mL PBS. Fluorescence intensity was measured using the Microplate Fluorometer (Varioskan Flash®, Thermo Scientific, Finland) at 485 nm and 498 nm excitation, as well as 530 nm and 585 nm emission for DCFH-DA and DHE assays, respectively.

**Analysis of cell viability**

T98G cell viability was determined using MTS assay according to the manufacturer’s protocol (Cell Titer 96Non-Radioactive Cell Proliferation Assay kit®, Promega, USA). Briefly, ten thousand cells after rotenone treatment were washed with PBS and added with 20 μl of MTS/ PMS mixture (MTS: PMS=20:1) to each well and incubated for 1–4 h under standard conditions (5% CO2 at 37°C). The quantity of formazan product was measured using spectrophotometer at 490 nm, and the absorbance was directly proportional to the number of living cells in culture.

**Analysis of cell proliferation**

T98G cell proliferation was analyzed using the Cell Proliferation ELISA, BrdU kit® (Roche, Germany) according to the manufacturer’s protocol. After rotenone treatment, 2×10^4 cells in each 24-well plate were labeled with 20 μL/well of BrdU and incubated for 24 h. The reaction product was quantified by measuring the absorbance at 490 nm using the Microplate Reader (Varioskan Flash®, Thermo Scientific, Finland). The developed color and thereby the absorbance values directly correlated to the amount of DNA synthesis and hereby to the number of proliferating cells.

**Analysis of cell apoptosis**

Cell apoptosis was analyzed using terminal deoxynucleotidyl transferase (TdT) dUTP Nick-End Labeling (TUNEL) assay (In Situ Cell Death Detection kit®, POD) according to the manufacturer’s protocol. This assay has been designed to detect apoptotic cells that undergo extensive DNA fragmentation during the late stages of apoptosis. Fifty thousand T98G cells were plated in slide chamber one day before rotenone treatment. The counterstaining process was performed using methyl green solution, as described previously [19]. Cells were observed under light microscope (Nikon ECLIPSE 80i). Dark brown cells were identified as apoptotic cells and calculated using Image J cell counter software (NIBHI).

**Analysis of MnSOD mRNA expression using real-time polymerase chain reaction (RT-PCR)**

Total RNA was isolated from harvested T98G cell culture using TriPure Isolation kit® (Roche) according to the manufacturer’s protocol. The concentration of total RNA was determined using spectrophotometry at 260 nm wavelength. cDNA of MnSOD was synthesized from 200 ng of total RNA samples and amplified using Taqman one-step RT-PCR Kit, according to the manufacturer’s protocol. 18S rRNA was used as a reference gene. The amplification protocol was performed using primers for MnSOD or 18S rRNA gene, as described in our previous report [20]. Each sample was performed in triplicate and the results were calculated using the 2^(-ΔΔCt) method.

**Analysis of MnSOD protein**

Total protein was isolated from harvested T98G cell culture using TriPure Isolation kit® (Roche) according to manufacturer’s protocol. Total protein concentration in each sample was first measured using spectrophotometer at 280 nm wavelength and plotted to the bovine serum albumin standard curve. The human MnSOD standard stock solution was reconstituted by adding the dilution buffer to obtain MnSOD concentration of 1600, 800, 400, 200, 100, 50, 25, and 0 pg/mL. MnSOD protein levels were determined using spectrophotometer at 450 nm and calculated using the concentration of standards provided in the kit and the total protein concentration in each sample.

**Analysis of the MnSOD enzyme-specific activity**

MnSOD enzyme activity was measured using xanthine oxidase inhibition assay (ranSOD® kit, Randox), as previously described [19]. To inhibit the Cu/ZnSOD first, nitric oxide (NO) was added into each sample, and the mixture was incubated for 5 min in room temperature [21]. MnSOD enzyme activity was expressed as percentage (%)/mg total protein.

**Analysis of transmission electron microscopy (TEM)**

The mitochondrial ultrastructure of T98G cells treated with rotenone was obtained by TEM. Cells were firstly fixed in 2.5% glutaraldehyde containing 3% sucrose in 0.1 M sodium cacodylate buffer (pH 7.4) for 24 h at 4°C, then in 2.5% glutaraldehyde containing 3% sucrose in 0.1 M sodium cacodylate buffer (pH 7.4) for 24 h at 4°C [22]. Afterward, samples were dehydrated in graded ethanol for 15 min, embedded in Spurr’s resin for 24 h at room temperature and examined under a JEOL 1010 transmission electron microscope.

**Statistical analysis**

Statistical analysis was performed using Student’s t-test (for comparison between rotenone- and DMSO-treated cells) or Wilcoxon test (for comparison of non-parametric data). Data were presented as mean ± SE, and p values of <0.05 were considered as being statistically significant.

**RESULTS**

Generation of intracellular ROS by rotenone in T98G cells

To determine the optimal rotenone concentration which could induce the overaccumulation of both superoxide and peroxide radicals intracellular, T98G cells were first treated with various concentrations of rotenone (0.5 μM, 5 μM and 50 μM) for 6 h. Fig. 1 demonstrates that rotenone could induce the production of both superoxide radicals.
Effect of rotenone treatment on cell survival

In this study, we investigated the effect of rotenone treatment on the survival of T98G cells. To analyze T98G cell viability following rotenone treatment, an MTS assay was performed. The result demonstrated that the viability of T98G cells treated with rotenone of either 5 or 50 µM was lower (~0.9 or 0.7-fold, p<0.05) than their counterparts treated with DMSO (Fig. 2a). These data were consistent with the superoxide level, revealing that superoxide radicals generated by rotenone were able to lessen T98G cell viability. To verify whether the decrease of T98G cell viability was triggered by the reduction of cell proliferation or induction of cell death, we performed BrdU assay for cell proliferation and TUNEL assay for cell apoptosis. The result demonstrated that T98G cells proliferation following rotenone treatment was slightly decreased (~0.9-fold; p>0.05) compared with the cells treated with DMSO (Fig. 2b).

In contrast to the result of cell proliferation, TUNEL assay in T98G cells treated with high-dose rotenone (50 µM) detected a remarkable increase of dark brown cells (~1.6-fold; p<0.05) compared with the control (non-treated cells) and DMSO-treated cells, respectively (Fig. 3). This indicates that high-dose rotenone provoked cell apoptosis which was most likely caused by overproduction of both superoxide and peroxide radicals.

Effect of rotenone treatment on MnSOD expression

To determine whether high-dose rotenone was efficient in stimulating the antioxidant response against the overproduction of ROS, we assessed the expression of MnSOD - a major antioxidant enzyme located in mitochondria - by gathering its level of mRNA and protein expression as well as its specific activity levels. In Fig. 4a and b, we found that the relative expression level of MnSOD mRNA in 50 µM rotenone-treated T98G cells was significantly upregulated (~3.8-fold; p<0.01). In contrast to the mRNA expression, high-dose rotenone led to a remarkable decrease in the protein levels (~6.2-fold lower; p<0.01) and specific activity of MnSOD in T98G cells (~1.4-fold; p<0.05). This revealed that high-dose rotenone modulated the MnSOD expression in T98G cells, leading to the modulations in its mRNA, protein, and activity levels.

DISCUSSION

Rotenone is an herbal pesticide that blocks electron flow from complex I to coenzyme Q of the electron transport chain in mitochondria [23]. As a lipophilic molecule, rotenone can freely penetrate through the cell membrane and coenzyme Q of the electron transport chain - could disrupt the mitochondrial integrity of T98G cells, leading to the modulations in its mRNA, protein, and activity levels.

Effect of rotenone treatment on the mitochondrial structure

To evaluate whether rotenone - an inhibitor of the mitochondrial transport electron chain - could disrupt the mitochondrial integrity of T98G cells, we observed the ultrastructure of mitochondria using TEM. The result illustrated that mitochondria of T98G cells treated with high-dose rotenone were swollen and exhibited irregular cristae with loss of lipid bilayer membrane (Fig. 5c), in contrast to the non-treated (Fig. 5a) and DMSO-treated cells (Fig. 5b), respectively. In the T98G cells treated with DMSO solely, although the mitochondria also tended to be swelling and irregular, the lipid bilayer membrane and cristae remained intact indicating that the mitochondria integrity was maintained (Fig. 5b). This observation demonstrated that rotenone treatment in T98G cells could lead to mitochondrial damage and cell death.

**Table 1: Fluorescence intensity of oxidized dichloro-dihydro-fluorescein diacetate and dihydroethidium in treated cells as expressed as ratio to control (cells without any treatment). All values are means, n=3. Student’s t-test showed significant differences at *(p<0.05) and **(p<0.01) compared to those of DMSO-treated cells. D: DMSO, R: Rotenone**

![Table 1](image1)

**Fig. 1: Effect of various rotenone concentrations on the reactive oxygen species (ROS) production. T98G cells were treated either with various concentrations of rotenone (0.5, 5, and 50 µM, respectively) in dimethyl sulfoxide (DMSO) or with the same volume of DMSO (as vehicle) used for dissolving rotenone. ROS levels were determined using DCFH-DA and DHE assay as described under "material and methods." Fluorescence intensity of oxidized dichloro-dihydro-fluorescein diacetate and dihydroethidium in treated cells was expressed as ratio to control (cells without any treatment). All values are means, n=3. Student’s t-test showed significant differences at *(p<0.05) and **(p<0.01) compared to those of DMSO-treated cells. D: DMSO, R: Rotenone.**

**Fig. 2: Effect of rotenone treatment on cell viability (a) and proliferation (b) on various concentration. T98G cells (1×10^5 cells) were first treated with 0.5 µM, 5 µM, and 50 µM rotenone in dimethyl sulfoxide (DMSO) or with DMSO (as vehicle) solely. Then, a MTS assay was performed for the cell viability and BrdU assay for cell proliferation, respectively, as described under the aforementioned material and methods section. Data were calculated as a percentage of the control (cells without any treatment). All values are means±standard error, n=9. Student’s t-test showed significant differences at *(p<0.05) and **(p<0.01) compared to the cells treated with DMSO.**

Wanandi et al.

membrane into the cytoplasm and mitochondria and easily transfer across the blood-brain barrier [23]. It has been reported that rotenone treatment could induce free radical generation, leading to oxidative damages, such as mitochondrial dysfunction, ubiquitin-dependent proteasome dysfunction and endoplasmic reticulum stress, as well as cell death [25,26]. In the present study, we confirmed that rotenone induced the accumulation of mitochondrial ROS, as reported previously [25,26]. Although superoxides were the main ROS generated as a result of mitochondrial complex I inhibition, here, we demonstrated that high-dose rotenone could stimulate the substantial production of peroxides as well. This might be due to the increased conversion of superoxides into hydrogen peroxides catalyzed by MnSOD in mitochondria.

The rotenone treatment of T98G cells performed in this study can serve as a model for oxidative stress induction during radiotherapy for glioma patients. Radiotherapy induces ROS production in tumor cells, either to facilitate tumor cell death through a process of apoptosis or to suppress the tumor growth through an inhibition of cell proliferation. Here, we found that rotenone concentration of 5 or 50 µM suppressed the viability of T98G cells, indicating high ROS levels [27]. The assessment of cell viability should be considered to be one of the primary criteria for apoptosis [28]. Through the use of a TUNEL assay, we observe an increase in DNA fragmentation which indicated cell death induced by chromatin dysfunction in T98G cells after high-dose rotenone treatment. Therefore, we suggested that the decline of cell viability following rotenone treatment is more likely due to the stimulation of cell apoptosis rather than the inhibition of cell proliferation since the BrdU assay performed in this study has indicated no significant decrease of T98G cell proliferation. Comparable results have also been reported by other previous studies, suggesting that rotenone does not affect the S-phase of cell proliferation [29,30]. Meanwhile, low-dose rotenone treatment (0.5-1 µM) for 18 h has been demonstrated to enhance apoptosis in HL-60 cells through cytochrome c release, caspase-3 activation, and DNA breakdown [26].

However, it should be considered that ROS-mediated DNA fragmentation presented by the TUNEL assay could be detected not only merely in apoptosis but also in the necrosis process [31]. In addition, using a TEM assay, we could observe the mitochondrial swelling with loss of cristae structure and loss of membrane integrity in the cells treated with high-dose rotenone, which is more likely an indicator of necrosis rather than apoptotic cell death, as described previously [32,33]. It has been suggested that an inhibitor of the mitochondrial electron transport chain such as rotenone-induced necrosis rather than apoptosis [33]. This effect was surely unexpected and should be taken into account during the management of cancer therapies in the future.

Fig. 3: Effect of rotenone treatment on DNA fragmentation. (a) T98G cells (1×10^5 cells) were first treated with various concentrations of rotenone in dimethyl sulfoxide (DMSO) and with an equivalent volume of DMSO (as vehicle), respectively. Subsequently, TUNEL assays were performed as described under the material and methods sectopm. Data were calculated as a percentage of control (cells without any treatment). All values are means±standard error, n=9. Student’s t-test showed significant differences at **(p<0.01) compared to the cells treated with DMSO. (b-i) micrographs of TUNEL-stained T98G treated with rotenone. Cells were observed under inverted microscope with ×200 magnification. (b) Negative control (cells without treatment); (c) positive control (cells + DNase); (d) cells + DMSO for 0.5 µM rotenone (vehicle); (e) cells + 0.5 µM rotenone; (f) cells + DMSO for 5 µM rotenone (vehicle); (g) cells + 5 µM rotenone; (h) Cells + DMSO for 50 µM rotenone (vehicle); (i) cells +50 µM rotenone. Apoptotic cells exhibiting DNA fragmentation were stained dark brown. Slides were counter-stained with methyl green to identify background TUNEL-negative cells.
higher than in benign tumor and normal tissue [12]. That MnSOD protein expression in ovarian carcinoma was significantly lower compared to normal brain cells [7]. In contrast to those results, mRNA synthesis was upregulated in high-grade GBM cells isolated resistance [40-42]. Indeed, our recent report has found that MnSOD has proposed that MnSOD acts as an anti-apoptotic factor. Thus, upregulation of MnSOD gene expression has been suggested as a specific properties of tumor cells [12,35-38]. In primary GBM multiforme, cancers which suggest that it may play a role in increasing the invasive initial thought to have tumor suppressor activity, but the recent studies free radical scavenging enzyme in the mitochondrial matrix. MnSOD was accumulation, it will induce a state of oxidative stress which inflicts variance oxidative damage on essential macromolecules in the cells, such as lipid, protein, and DNA [34]. Therefore, we focused to analyze the response of rotenone induction on the expression of MnSOD, the major free radical scavenging enzyme in the mitochondrial matrix. MnSOD was initially thought to have tumor suppressor activity, but the recent studies have proved that the expression of this enzyme was upregulated in many cancers which suggest that it may play a role in increasing the invasive properties of tumor cells [12,35-38]. In primary GBM multiforme, upregulation of MnSOD gene expression has been suggested as a specific diagnostic marker through transcriptome analysis [39]. Several studies have proposed that MnSOD acts as an anti-apoptotic factor. Thus, upregulation of MnSOD gene expression could lead to ROS-based therapy resistance [40-42]. Indeed, our recent report has found that MnSOD mRNA synthesis was upregulated in high-grade GBM cells isolated from clinical specimens, but its protein and specific activity levels were lower compared to normal brain cells [7]. In contrast to those results, other previous study that used tissue microarray analysis has shown that MnSOD protein expression in ovarian carcinoma was significantly higher than in benign tumor and normal tissue [12].

Interestingly, the present study hinted that both the protein level and specific activity of MnSOD in the cells treated with high-dose rotenone were significantly lower than those in control cells, which contradicted findings in the previous reports [12,26]. In addition, we specifically employed the analysis of MnSOD enzyme-specific activity with a xanthine oxidase inhibition assay and inhibited Cu/ZnSOD activity by adding natrium cyanide [20]. It should be noted that the rotenone dose optimized in this study was higher than the dosages used in the aforementioned studies and presuming that the overaccumulation of intracellular ROS may exceed the antioxidant capacity of MnSOD, which reduces the availability and activity of MnSOD. More interestingly, the present study revealed that high-dose rotenone (50 µM) could modulate mRNA, protein, or specific activity levels of MnSOD in T98G cells. Although the synthesis of MnSOD mRNA has been upregulated, accumulation of intracellular ROS was elevated, as indicated by high superoxide and peroxide levels, which led to insufficient levels of active intracellular MnSOD enzyme. Thus, the imbalance between high ROS level and low activity of MnSOD found in the present study was due to high-dose rotenone-induced oxidative stress.

On the other hand, we should also consider the possibility that ubiquitin-proteasomal degradation of MnSOD protein is exaggerated by the high abundance of intrinsic ROS leading to mitochondrial dysfunction [25,26]. This assumption is supported by the TEM assay results in the present study demonstrating that the mitochondrial disruption in T98G cells is induced by high ROS levels. Alternatively, the discrepancy between high MnSOD mRNA and low protein or specific activity level might be associated with the dysregulation of MnSOD protein synthesis influenced by overproduction of ROS. The steady state of mRNA levels perceived using quantitative RT-PCR method is a cumulative result of several regulatory mechanisms, including transcription, RNA processing, and RNA stability, whereas the protein level and activity are predominantly influenced by translational and post-translational processing as well as innate variations in stability. Several comparative transcriptomics and proteomics in mouse offer evidence that there is a low correlation between transcript and encoded protein levels [43,44].

Here, we highlight the importance of the MnSOD gene expression analysis at the level of mRNA and protein, as well as enzyme activity. Unlike the previous report on ovarian cancer [12], our concomitant study has obtained that the suppression of MnSOD mRNA expression in T98G cells through siRNA transfection could reduce the protein level and enzyme activity and enhance superoxide radicals production, leading to cell death [45]. Hence, we evidently suggest that the impact of oxidative stress induced by high-dose rotenone is most likely associated with low MnSOD expression at the level of protein and enzyme activity. Increased
MnSOD mRNA levels which resulted in the increased availability of MnSOD antioxidants might be the result of a cellular adaptive response to oxidative stress. It should be considered that the impact of rotenone on cell survival is not only limited through enhancing the amount of intracellular ROS, both superoxide and peroxide radicals, but also through the modulation of MnSOD gene expression.

Based on the results of this study, we could elucidate that the discrepancy between high MnSOD mRNA and low protein or specific activity levels in human high-grade glioma cells reported in our previous study might be due to oxidative damage triggered by excessive production of ROS, as demonstrated by the increase of protein carbonyls as a marker of protein oxidative damage in the high-grade compared to low-grade glioma and normal brain cells [7]. Nevertheless, in contrast to the present study, the rate of cell proliferation was higher, and the rate of apoptosis was lower in the high-grade compared to low-grade glioma cells. It seems that the redox signaling pathway regulating cell viability was not similar to that observed in this study. It has been formerly reported that moderate ROS levels were able to induce cell proliferation through a redox signaling pathway, which contributed to tumor growth, whereas high ROS levels stimulated apoptosis [27]. Furthermore, in that study, we have demonstrated that cell survival of high-grade glioma from clinical specimens was strongly correlated with the low MnSOD mRNA level [7]. Therefore, we presume that MnSOD gene expression might be responsible for the modulation of redox signaling mechanisms contributing to tumor cell survival in high-grade glioma cells, and further affecting tumor malignancy and ROS-based cancer therapy resistance.

CONCLUSION

Based on our findings, we conclude that overproduction of ROS in rotenone-treated human GBM T98G cells could suppress the MnSOD protein level and activity even though its mRNA synthesis has increased. This diminished cell survival rates through enhancement of cell death rather than inhibition of cell proliferation. Suppression of MnSOD protein level and activity might be beneficial for radiotherapy. However, the upregulation of MnSOD mRNA synthesis during ROS-based cancer therapy should be considered, since it would maintain the availability of MnSOD activity and may also reestablish cell viability and proliferation. Therefore, the suppression of MnSOD synthesis in glioma cells before radiotherapy could be proposed as a targeted therapy to treat radiation resistance of GBM.

ACKNOWLEDGMENT

This research was funded by Riset Unggul Universitas Indonesia (RUUI) 2010. The authors would like to express our gratitude to Direktorat Riset dan Pengabdian Masyarakat Universitas Indonesia for writing assistances.

REFERENCES


30. Wang XJ, Xu JX. Possible involvement of Ca2+ signaling in