Adriamycin-induced, TNF-α-mediated central nervous system toxicity

Jitbanjong Tangpong, a,b Marsha P. Cole, a Rukhsana Sultana, c Gururaj Joshi, c Steven Estus, d Mary Vore, a William St. Clair, c Suvina Ratanachaiyavong, b Daret K. St. Clair, a and D. Allan Butterfield c,*

a Graduate Center for Toxicology, University of Kentucky, Lexington, KY 40536, USA
b Faculty of Medicine, Prince of Songkla University, Songkhla, Thailand
c Department of Chemistry, Center of Membrane Sciences, and Sanders-Brown Center on Aging, 121 Chemistry-Physics Building, University of Kentucky, Lexington, KY 40506, USA
d Department of Physiology, University of Kentucky, Lexington, KY 40536, USA

e Department of Radiation Medicine, University of Kentucky, Lexington, KY 40536, USA

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The clinical effectiveness of adriamycin (ADR), a potent chemotherapeutic, is known to be limited by severe cardiotoxic side effects. However, the effect of ADR on brain tissue is not well understood. It is generally thought that ADR is not toxic to the brain because ADR does not pass the blood–brain barrier. The present study demonstrates that ADR autofluorescence was detected only in areas of the brain located outside the blood–brain barrier, but a strong tumor necrosis factor (TNF) alpha immunoreactivity was detected in the cortex and hippocampus of ADR-treated mice. Systemic injection of ADR led to a decline in brain mitochondrial respiration via complex I substrate shortly after ADR treatment (P < 0.05). Cytochrome c release, increased caspase 3 activity, and TUNEL-positive cell death all were suggestive of apoptosis in brain following systemic ADR treatment. The levels of the known pro-apoptotic proteins, p53 and Bax, were increased in brain mitochondria at 3 h following ADR treatment and declined by 48 h. In contrast, the anti-apoptotic protein, Bcl-xL, was increased later at 6 h post-ADR treatment and was sustained throughout 72 h. Furthermore, p53 migrated to mitochondria and interacted with Bcl-xL, supporting the hypothesis that mitochondria are targets of ADR-induced CNS injury. Neutralizing antibodies against circulating TNF completely abolished both the increased TNF in the brain and the observed mitochondrial injury in brain tissues. These results are consistent with the notion that TNF is an important mediator by which ADR induces central nervous system (CNS) injury. This study, the first to provide direct biochemical evidence of ADR toxicity to the brain, revealed novel mechanisms of ADR-induced CNS injury and suggests a potential therapeutic intervention against circulating TNF-induced CNS effects. © 2006 Elsevier Inc. All rights reserved.

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Introduction

Adriamycin (ADR), an antibiotic produced by the fungus Streptomyces peucetius, is a potent anticancer drug commonly used in the treatment of a variety of cancers including breast cancer (Hitchcock-Bryan et al., 1986; Fisher et al., 1989). However, its clinical effectiveness is limited by the toxic effect on normal tissues (Singal et al., 1987, 2000; Meredith and Reed, 1983; Oteki et al., 2005), including a cumulative, dose-related cardiomyopathy (Singal and Iliskovic, 1998). Recent studies in breast cancer survivors have shown persistent changes in cognitive function, including memory loss, tendency for distractions, and difficulty in performing multiple tasks, following chemotherapy (Schagen et al., 1999; Brezden et al., 2000). These studies report that cognitive deficits, particularly in the areas of memory and concentration, are associated with cancer chemotherapy regimens, both in the short-term after treatment, and up to 2 years and more than 5 years after diagnosis (Ahles et al., 2002; Ferrell and Hassay Dow, 1997). These cognitive problems, collectively called somnolence or cognitive dysfunction, are also reported in cancer patients undergoing adriamycin-based chemotherapy, especially breast cancer patients (Freeman and Broshek, 2002; Schagen et al., 2001; Meyers, 2000).

Although the biochemical basis for these cognitive problems is unknown, it has been demonstrated that cancer therapeutic agents
such as ADR can modulate endogenous levels of cytokines such as tumor necrosis factor (TNF) alpha (Usta et al., 2004). Enhanced circulating TNF can initiate local TNF production via activation of glia cells leading to production of reactive oxygen/nitrogen species (RONS) (Szelenyi, 2001). RONS, including superoxide, hydrogen peroxide, and nitric oxide, can react directly with each other or indirectly to generate even more reactive species (Halliwell and Gutteridge, 1999). We recently reported that 72 h after a single i.p. injection of ADR, there was a significant increase in levels of protein oxidation and lipid peroxidation in brain tissues (Joshi et al., 2005). However, the mechanism by which ADR causes oxidative stress in the brain remains unknown.

It is well established that ADR does not cross the blood–brain barrier (Bigotte and Olsson, 1982; Bigotte et al., 1982), but that circulating levels of TNF can directly pass the blood–brain barrier and activate microglia and neurons to further increase local TNF levels (Osburg et al., 2002). TNF is known to induce neuronal damage (Gutierrez et al., 1993). TNF-induced tissue injury is mediated, at least in part, by its effect on mitochondria (Goossens et al., 1995). TNF induces morphologic damage of mitochondria and biochemical respiratory defects in cultured cells (Liu et al., 2004; Schulz-Osthoff et al., 1992). The cytotoxicity of TNF depends on the induction of the mitochondrial permeability transition pore (Lancaster et al., 1989). Thus, it is possible that an increase in TNF levels may be a link between ADR-induced oxidative stress and CNS injury.

The present study evaluated the relationship between ADR-induced TNF production, mitochondrial dysfunction, and CNS injury. The results provide biochemical insights into the mechanisms of ADR-induced CNS injury.

Materials and methods

Animals

Eight-week-old male B6C3 mice (25–30 g) were kept under standard conditions, and all experimental procedures were approved by the Institutional Animal Care and Use Committee of the University of Kentucky.

Treatments

Mice were injected in a single intraperitoneal (i.p.) dose of 20 mg/kg adriamycin (doxorubicin hydrochloride, Gensia Sicor Pharmaceuticals, Inc., Irvine, CA) or the same volume of saline as control for 3 h. This dose and time were based on previous studies in which we demonstrated ADR-induced cardiomyopathy (Yen et al., 1996, 1999).

To determine whether the neutralization of TNF in the periphery would mitigate the brain biochemical effects of ADR, anti-mouse TNF antibody (R&D Systems, Minneapolis, MN) was diluted in saline and immediately injected in a single i.p. dose of 40 ng/kg, and anti-TNF antibody immediately followed by ADR also was injected to separate animals. Controls consisted of anti-TNF antibody alone or preimmune rabbit IgG or saline in the same total volume. All results were obtained from at least three separate experiments.

Localization of adriamycin in brain tissues

Mice were euthanized by the i.p. injection of 65 mg/kg of Nembutal (Sodium pentobarbital, Abbott Laboratories, North Chicago, IL) and were perfused via cardiac puncture initially with 0.1 M phosphate-buffered saline (PBS), pH 7.4, and subsequent fixation with 4% paraformaldehyde. Brain tissues were removed and coronal cryosection at regular intervals of 7-μm thickness was performed. The sections were prepared for immunohistochemistry and detection of ADR. ADR in brain tissue slices was directly visualized using an inverted fluorescence microscope (excitation filter 550 nm, and barrier filter 590 nm). Photomicrographs were taken with an Olympus MagnaFire digital camera (Olympus, America, Melville, NY).

Enzyme-linked immunosorbent assay (ELISA)

Mice were treated with 20 mg/kg ADR or saline as control. Blood samples were collected at 1, 3, 6, 9, and 24 h and allowed to clot at 2–8°C overnight. Serum samples were used to measure TNF levels, according to the mouse enzyme-linked immunosorbent assay following the manufacturer’s instructions (mouse TNF-α/ TNFSF1A immunoassay, R&D Systems, Minneapolis, MN). The TNF concentration in the sample was calculated from a recombinant mouse TNF standard curve. The minimum detection limit is typically less than 5.1 pg/ml.

Immunohistochemistry study

Brain tissue slices were fixed in 4% paraformaldehyde for 15 min, air dried, and washed with PBS. Nonspecific proteins were blocked in blocking serum, consisting of 3% normal donkey’s serum, and 0.3% Triton X-100 in PBS, and incubated at room temperature for 30 min. After blocking, slices were incubated with primary anti-TNF (Upstate, Lake Placid, New York). Anti-MAP2 antibody (Chemicon, Temecula, CA) was used as a neuronal marker in order to study the location of TNF in neurons of cortical and hippocampal regions. The sections were kept in a humidified box at 4°C overnight. Tissues were washed three times with PBS and then were incubated for 1 h with donkey antibody-conjugated secondary antibodies conjugated with fluorescent dyes. Excess secondary antibodies were removed by washing three times in PBS and once with deionized H2O. Tissue slides were mounted with mounting medium (Vectorshield, H-100, Vector Laboratories, Burlingame, CA). Photomicrographs were obtained using a Leica confocal fluorescence microscope (Leica Microsystems Inc., Bannockburn, IL, USA).

Mitochondrial isolation and purification

Mice were perfused via cardiac puncture with cold mitochondrial isolation buffer, the brain promptly removed, the cerebellum dissected away, and the mitochondria immediately isolated from the brain by a modification of the method described by Mattiazzi et al. (2002). Brain mitochondria were isolated in cold mitochondrial isolation buffer, containing 0.07 M sucrose, 0.22 M mannitol, 20 mM HEPES, 1 mM EGTA, and 1% bovine serum albumin, pH 7.2. Tissues were homogenized with a Dounce homogenizer and centrifuged at 1500 × g at 4°C for 5 min before transferring the supernatants. The pellets were resuspended and centrifuged at 1500 × g at 4°C for 5 min. The supernatants were combined and recentrifuged at 1500 × g at 4°C for 5 min. The supernatants were separated and centrifuged at 13,500 × g at 4°C for 10 min. Mitochondrial pellets were resuspended in 50–100 μL cold mitochondrial isolation buffer. Protein concentration of isolated
Mitochondria was determined by the Bradford assay (Bradford, 1976). The final protein concentration was 20–40 mg/ml.

Mitochondrial respiration

Mitochondrial respiration was determined using Clark-type polarographic oxygen sensors (Hansatech Instruments, Kings Lynn, Norfolk, UK) to measure the rate of oxygen consumption. Freshly isolated mitochondria were suspended in respiration buffer at a concentration of 0.5 mg mitochondrial protein per milliliter of respiration buffer, which consists of 0.25 M sucrose, 50 mM HEPES, 1 mM EGTA, 10 mM KH$_2$PO$_4$, and 2 mM MgCl$_2$, pH 7.4. Oxygen consumption was measured with either pyruvate (10 mM) plus malate (10 mM) or succinate (10 mM) as substrates for respiration from complex I or complex II in the absence of exogenous ADP (state II) and after addition of 300 mM ADP (state III respiration). Rotenone (5 $\mu$M) was added to the reaction to inhibit respiration from complex I when succinate was used as the substrate. The ATPase inhibitor oligomycin (100 $\mu$g/ml) was added to inhibit mitochondrial respiration such that state IV respiration was similar to the state II respiration rate. FCCP (1 $\mu$M), an uncoupling agent, was added as a control of respiration. Respiration control ratios (RCR) were calculated as the ratios of state III and state II respiration as described previously by Estabrook (1967). The unit for state II rate and state III rate is nmol/min/mg protein.

Preparation of brain homogenates

Brains perfused with PBS were isolated and dissected from six groups of mice 3 h postinjection of a single dose of ADR, anti-TNF antibody, anti-TNF antibody immediately followed by ADR, IgG, or saline-treated mice. Brain was isolated and placed in 0.1 M PBS, pH 7.4, containing protease inhibitors, 4 $\mu$g leupeptin, 4 $\mu$g pepstatin, and 5 $\mu$g aprotinin, washed and minced in ice-cold PBS containing...
protease inhibitors. Tissues were homogenized with a Dounce homogenizer and centrifuged at 12,500 × g at 4°C for 30 min before transferring the supernatant. Protein concentration of brain homogenate was determined by the Bradford assay (Bradford, 1976).

Western blot analysis

Perfused brain homogenates and isolated mitochondrial proteins were separated via 12.5% denaturing polyacrylamide gel electrophoresis (SDS–PAGE) and transferred to a nitrocellulose membrane. The membrane was blocked for 1 h at room temperature in blocking solution consisting of 5% nonfat dried milk, 0.5% Tween-20, and Tris-buffered saline (TBST), pH 7.9. After blocking, the membrane was incubated overnight at 4°C with primary antibodies against TNF (Upstate, Lake Placid, New York) and β-actin (Clone AC-74, A5316, Sigma, Saint Louis, MO) in homogenate samples. For isolated mitochondria samples, the membrane was incubated overnight at 4°C with primary

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Fig. 4. Immunofluorescence analysis of TNF localization in cortex following ADR treatment. Confocal microscopy analysis of TNF (red) and neurons (MAP2, green) and colocalization (yellow) showed that TNF is increased in neurons of mice 3 h after treatment with 20 mg/kg ADR compared to mice treated with saline (A), anti-TNF-antibody together with ADR, or anti-TNF antibody alone (B). Pictures are representative images from at least 3 independent experiments per group.
antibodies against p53 (Ab-11, Oncogene Research, Cambridge, MA), Bcl-xL (S-18), Bax (P-19), succinate dehydrogenase (SDHB) (Santa Cruz Biotechnology, Santa Cruz, CA), in blocking solution. The membrane was washed twice in TBST and incubated for 1 h with horseradish peroxidase-conjugated secondary antibodies in blocking solution. After incubation with secondary antibodies, the membrane was washed twice with TBST and once in TBS (TBS without 0.5% Tween-20). Immunoreactivities of the protein bands were detected by enhanced chemiluminescence autoradiography (ECL, Amersham Phamacia Biotech, Arlington Heights, IL) as described by the manufacturer.

To further investigate cytochrome c release, mitochondrial and cytosolic fractions were isolated from brain tissues and were size separated and probed with anti-cytochrome c antibody (BD Biosciences, CA). Succinate dehydrogenase (SDHB) and β-actin were used as loading controls employing the procedure described above.

Fig. 5. Immunofluorescence analysis of TNF localization in hippocampus following ADR treatment. Confocal microscopy analysis of TNF (red) and neurons (MAP2, green) and colocalization (yellow) showed that TNF is increased in neurons of mice 3 h after treatment with 20 mg/kg ADR compared to mice treated with saline (A), anti-TNF-antibody together with ADR, or anti-TNF antibody alone (B). Pictures are representative images from at least 3 independent experiments per group.
Immunoprecipitation assays

Isolated mitochondrial proteins (500 μg) were resuspended in 500 μL RIPA buffer (9.1 mM Na₂HPO₄, 1.7 mM NaH₂PO₄, 150 mM NaCl, 0.5% sodium deoxycholate, 1% v/v Nonidet P40, 0.1% sodium dodecyl sulfate, pH 7.2). Protease inhibitors (0.1 mg PMSF and 1 μg aprotinin per ml RIPA) were added at the time of use followed by incubation with 5 μg/ml of mouse p53 antibody (Ab-11, Oncogene) at 4°C overnight. Protein A/G-Agarose (50 μL) was added to the reaction mixture with the antibody. Immunocomplexes were collected by centrifugation at 1000 g at 4°C for 5 min and then washed four times with RIPA buffer. Immunoprecipitated samples were recovered by resuspending in 2× sample loading buffer, and Bcl-xL (Santa Cruz Biotechnology, Santa Cruz, CA) proteins were detected by Western blot. In employing IgG, preimmune serum was used in procedures to immunoprecipitate the isolated mitochondrial proteins and subsequent analysis of the pellet and supernatant by Western blot using an antibody against Bcl-xL in order to exclude the interference from light chain. Immunoreactivity was evaluated on immunoblots by densitometric analysis using a Bio-RAD densitometer (Bio-RAD Laboratory, Inc, Hercules, CA, USA).

Caspase 3 activity assay

Caspase 3 activity assay was performed with the use of a colorimetric substrate in accordance with the protocol supplied by the manufacturer (Stigma, St. Louis, MO). In brief, mice were anesthetized and perfused with 1× PBS to reduce any enzyme activity associated with intravascular blood components. Brain was dissected and homogenized in lysis buffer containing 50 mM HEPES, pH 7.4, 5 mM CHAPS, 5 mM DTT plus 1 μg/ml aprotinin and pepstatin. The brain homogenates were incubated on ice for 15 min and then centrifuged at 16,000 g and then washed four times with RIPA buffer. Immunoprecipitated samples were recovered by resuspending in 2× sample loading buffer, and Bel-xL (Santa Cruz Biotechnology, Santa Cruz, CA) proteins were detected by Western blot. In employing IgG, preimmune serum was used in procedures to immunoprecipitate the isolated mitochondrial proteins and subsequent analysis of the pellet and supernatant by Western blot using an antibody against Bel-xL in order to exclude the interference from light chain. Immunoreactivity was evaluated on immunoblots by densitometric analysis using a Bio-RAD densitometer (Bio-RAD Laboratory, Inc, Hercules, CA, USA).

TUNEL assay

The assay was performed following the manufacturer’s instructions (Promega, Madison, WI). Briefly, the cryosections of brain tissues were fixed with 4% paraformaldehyde, permeabilized with Triton X-100, and incubated with biotinylated deoxynucleotidyltransferase (rTdT) for 1 h at 37°C. The fragmented DNA labeled at terminal transferase. Negative controls consisted of specimens in which deoxynucleotidyltransferase was omitted.

Statistical analysis

Statistical comparisons were made using one-way ANOVA followed by Newman–Keuls multiple comparisons test. Data are expressed as mean ± SEM.

Results

Adriamycin accumulation in brain tissues

To explore the possibility that ADR accumulates in brain tissue, mice were given 20 mg/kg ADR by a single i.p. injection. ADR accumulation in the CNS was studied by direct ADR fluorescence in brain slices using an inverted fluorescence microscope. The specific orange-red fluorescence of ADR was observed in several areas outside the blood–brain barrier, including the choroid plexus as previously reported by Bigotte and Olsson (1982) and Bigotte et al. (1982). ADR fluorescence was clearly distinguishable from the background of untreated control but was not observed in cortex and hippocampus (Fig. 1). When anti-TNF antibody and ADR were injected into mice together, ADR fluorescence was still observed (Fig. 1B). Fluorescence was not observed in mice treated with anti-TNF antibody alone (Fig. 1C).

Adriamycin-induced circulating TNF levels

TNF levels in serum were significantly higher in mice treated with ADR than those from controls (*P < 0.001; Fig. 2). The increased TNF levels were detectable as early as 1 h and were sustained throughout a 24-h period after ADR treatment.

Increased TNF level in brain tissues

TNF levels were significantly increased in brain tissue homogenates as detected by Western blot analysis, following administration of ADR (Fig. 3). Anti-TNF antibody blocked ADR-mediated increased brain levels of TNF (Fig. 3). However,
nonspecific IgG was unable to block ADR-induced TNF elevation in brain (Fig. 3). To determine the localization of TNF in brain tissues, tissue slices were stained with anti-TNF and anti-MAP2 antibodies to localize TNF. The TNF levels clearly were increased in neurons of cortex and hippocampus compared with saline controls because MAP2 is a neuron-specific marker (Figs. 4 and 5). To verify that the observed TNF in the brain was mediated by ADR-induced circulating TNF, we co-injected a neutralizing antibody against TNF along with ADR. The increased levels of TNF in cortical and hippocampal regions or in whole brain homogenates were blocked in mice treated with anti-TNF antibody and ADR (Figs. 3–5).

**Adriamycin-induced mitochondrial dysfunction**

To investigate the effect of ADR on brain mitochondrial function, brain mitochondrial respiration using pyruvate plus malate and succinate as the substrates was measured. The data are presented as the RCR of each treatment group and the IgG or saline-treated control group from each set of experiments. As shown in Fig. 6, the values of RCR from state III and state II respiration, via complex I but not complex II (data were not shown), were significantly decreased in the ADR or IgG followed by ADR treatment groups compared to controls (*P < 0.05). In mice treated with anti-TNF antibody only, or in mice treated with anti-TNF antibody followed by ADR, brain mitochondrial respiration was not significantly different from that of the control group. Taken together, these results indicate that ADR-induced circulating TNF levels subsequently increased brain levels of TNF, which led to inhibition of the NAD-linked state III respiration rate. The latter is mediated through complex I but not complex II. The anti-TNF antibody prevented the decline in mitochondrial respiration of brain tissues, consistent with this notion.

**Pro-survival and pro-apoptotic protein levels in mitochondria**

To probe the possibility that ADR-induced increased levels of serum and brain TNF are associated with altered levels of pro- and anti-apoptotic proteins in brain mitochondria, the levels of the pro-apoptotic proteins, p53 and Bax, and the anti-apoptotic proteins, Bcl-xL, were quantified (Fig. 7). The results demonstrate that p53 was increased in mitochondria at the earliest time point examined (3 h after ADR treatment compared with saline control (*P < 0.01). This elevated p53 level persisted at 6 and 24 h, but then declined at 48 and 72 h. Bax increased with a similar kinetics to that of p53 (*P < 0.01). The anti-apoptotic protein Bcl-xL increased after treatment with ADR at 6, 24, 48, and 72 h compared with saline control (*P < 0.05). The level of succinate dehydrogenase was not changed and was used for a normalization protein loading control. These results demonstrated a rapid increase of pro-apoptotic proteins and anti-apoptotic Bcl-xL in brain mitochondria of ADR-treated mice, which are associated with increased neuronal TNF. Consistent with the results shown above, blocking ADR-mediated elevated TNF with anti-TNF antibody resulted in no

![Fig. 7. Representative immunobots showing the levels of p53, Bax, Bcl-xL, and succinate dehydrogenase in mitochondria. Mitochondrial proteins were isolated from brain tissues of ADR- and saline-treated mice and separated by SDS–polyacrylamide gel electrophoresis. The pro-apoptotic protein p53 was significantly increased at 3, 6, and 24 h (*P < 0.01), and the pro-apoptotic protein Bax was increased at 3 and 6 h (*P < 0.01). The anti-apoptotic protein, Bcl-xL, showed an increased protein density 6, 24, 48, and 72 h after treatment with ADR (*P < 0.05). Succinate dehydrogenase was used to normalize protein loading. The results shown are a representative independent set of data of n = 3 separate sets from individual animals.](image-url)
increase of pro-apoptotic proteins in mitochondria 3 h after treatment with ADR (Fig. 8).

**p53 forms specific complex with the protective Bcl-xL protein**

p53 can participate in induction of apoptosis by acting directly at the mitochondria (Marchenko et al., 2000; Mihara et al., 2003). Localization of p53 to mitochondria occurs in response to apoptotic signals and precedes cytochrome c release and procaspase-3 activation (Schuler and Green, 2001). To determine the ability of p53 to interact with Bcl-xL in brain mitochondria, immunoprecipitation was performed using an antibody to p53 to precipitate mitochondrial proteins, and the complexes were probed with antibodies to p53 and Bcl-xL by Western blot analysis. The results showed specific increases of p53 and Bcl-xL in the ADR treatment groups compared to the controls (Fig. 9A; *P < 0.05). Immediately prior anti-TNF antibody treatment blocked p53 and BAX translocation to mitochondria following ADR treatment. Succinate dehydrogenase was used to normalize for protein loading. These results shown are a representative set of data of n = 3 separate sets from individual animals.

**Cytochrome c release from the mitochondria to cytosol**

We also examined whether ADR treatment led to brain mitochondrial membrane pore opening as assessed by cytochrome c release. Cytochrome c release from mitochondria and elevation in cytosol were found in the brain of mice treated with ADR or IgG followed by ADR compared with brain mitochondria isolated from mice treated with saline or IgG alone (*P < 0.01). Anti-TNF antibody prevented cytochrome c release from mitochondria to cytosol (Fig. 10).

**Increased caspase 3 activity and apoptosis**

Release of cytochrome c from mitochondria to cytosol induces caspase 3 cleavage and apoptosis in brain tissues. Three hours following treatment of mice with ADR, increased caspase 3 activity was observed in brain tissues (*P < 0.001). There was a progressive increase in caspase 3 activity after ADR treatment for 72 h (*P < 0.001; Fig. 11). TUNEL staining demonstrated dark brown nuclear condensation characteristic of apoptotic cell death corresponding to the time point of elevated caspase 3 activity in ADR-treated mice compared with saline control (*P < 0.01). DNA damage was greater at 72 h compared with 3 h in brain from ADR-treated mice (**P < 0.05; Fig. 12).

**Discussion**

A somnolence syndrome (also known as cognitive dysfunction), which often is called “chemobrain” by cancer patients receiving ADR (Wefel et al., 2004), has been unexplored, possibly due to the accepted notion that ADR does not pass the blood–brain barrier. In this report, we confirmed that ADR accumulated only in areas outside the blood–brain barrier but increased TNF
levels were found in serum and both the hippocampal and cortical regions of the brain. Mitochondrial function was altered in brain following ADR treatment. Furthermore, a neutralizing antibody against TNF given systemically abolished the observed TNF levels in brain tissue. Our results support the hypothesis that TNF is an important mediator of the observed ADR-induced mitochondrial dysfunction in brain.

Systemic TNF is well recognized to act as a signal in the complex network of immune–neuron interaction, which affects the CNS (Besedovsky and del Ray, 1996; Blattleis and Sehic, 1998; Licinio and Wong, 1997), such as in HIV and Alzheimer’s disease (Eric et al., 2002; Mattson et al., 2005; Valcour et al., 2004; Greig et al., 2004; Pocernich et al., 2005). Our finding that the tissue levels of TNF increase in brain tissue is consistent with the effect of ADR on circulating TNF levels. Circulating TNF may enter the brain by transport across the blood–brain barrier and stimulation of local TNF production after physically entering the brain (Gutierrez et al., 1993; Osburg et al., 2002). Alternatively, increased brain TNF levels may result from the activation of microglia and macrophages that enter the brain after ADR treatment. TNF can act on brain cells to cause the observed decline in mitochondrial respiration and subsequent increase in oxidative stress markers (Joshi et al., 2005). This possibility is strongly supported by our finding that neutralizing antibody against TNF successfully alleviated the decline in mitochondrial respiration of brain tissue after treatment with ADR. Our results are consistent with those previously reported by Usta et al. (2004), who showed that pentoxiphylline (PTX), an inhibitor of TNF-α, prevented an ADR-mediated systemic increase in TNF and nephropathy.

Fig. 9. Representative co-immunoprecipitation of the anti-apoptotic protein, Bcl-xL, in mitochondria. (A) Brain mitochondria were isolated from mice after ADR or saline (control) treatment and were immunoprecipitated with anti-p53 antibody. Bands specific for Bcl-xL were increased at 3 h (*P < 0.05). (B) Blocking with anti-TNF antibody prevented p53 translocation to mitochondria and co-localization with Bcl-xL in mice 3 h after injection with anti-TNF antibody immediately followed by ADR compared to mice treated with ADR or IgG followed by ADR (*P < 0.05). Similar immunoprecipitation studies employing IgG using preimmune serum revealed no band of Bcl-xL in the mitochondrial pellet fraction. The results shown are a representative set of data of n = 3 separate sets from individual animals.
Our results indicate that ADR induced a decline in brain mitochondrial respiration complex I but had no effect on complex II. This is consistent with the possibility that the [4Fe–4S] cluster protein in complex I is inactivated by ROS generated in mitochondria because the [4Fe–4S] protein of complex I extends into the inner membrane where superoxide is generated. This possibility is strongly supported by our previous studies, which demonstrated that transgenic mice overexpressing MnSOD are protected from ADR-induced complex I inactivation in cardiac tissues (Yen et al., 1996, 1999). Mitochondrial dysfunction may be one of the signals initiating the mitochondrial apoptosis pathway by translocation of pro-apoptotic proteins, p53 and Bax, to mitochondria, which in turn led to cytochrome c release and subsequent induction of caspase 3 cleavage and apoptotic cell death. The increase of these pro-apoptotic proteins coincides with the increase in mitochondrial dysfunction, suggesting mitochondrial dependent tissue injury. Our finding that p53 interacted with Bcl-xL in mitochondria further supports the role of mitochondria in ADR-induced complex I inactivation in cardiac tissues (Yen et al., 1996, 1999). Mitochondrial dysfunction may be one of the signals initiating the mitochondrial apoptosis pathway by translocation of pro-apoptotic proteins, p53 and Bax, to mitochondria, which in turn led to cytochrome c release and subsequent induction of caspase 3 cleavage and apoptotic cell death. The increase of these pro-apoptotic proteins coincides with the increase in mitochondrial dysfunction, suggesting mitochondrial dependent tissue injury. Our finding that p53 interacted with Bcl-xL in mitochondria further supports the role of mitochondria in ADR-induced complex I inactivation in cardiac tissues (Yen et al., 1996, 1999). 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resulting in cytochrome c release (Mihara et al., 2003). Over-expression of the anti-apoptotic protein Bcl-xL abrogates stress signal-mediated mitochondrial p53 accumulation and apoptosis (Marchenko et al., 2000).

Our results support the notion that p53 and Bax are translocated to mitochondria early following ADR treatment, which is the same kinetics for mitochondrial membrane permeabilization and cytochrome c release-initiated apoptotic cell death. The stabilization of p53 by its interaction with Bax prevents p53 degradation by the metalloprotease MDM2 (Tan et al., 2001). In response to this pro-apoptotic stress, our results suggest that the brain tries to compensate to promote cell survival by increased induction of the anti-apoptotic protein Bcl-xL, which is increased later at 6 h following ADR treatment.

In summary, our results for the first time provide direct biochemical evidence of ADR toxicity to brain. In particular, our results demonstrate that ADR-induced circulating TNF is causally related to the observed CNS injury associated with this cancer chemotherapy agent. TNF-induced mitochondrial dysfunction with its downstream consequences leading to further increases in oxidative stress in the brain may, at least in part, be responsible for the cognitive dysfunction (somnolence syndrome) observed in many patients undergoing ADR-based chemotherapy. Whether ADR-induced circulating TNF is directly responsible for the observed brain mitochondrial dysfunction or circulating TNF further increased TNF from activated glia cells remains to be determined but merits further investigation. Our results, which demonstrated that a neutralizing antibody against TNF abolished the observed mitochondrial injury in animals treated with ADR, are highly encouraging because antibodies against TNF have been in clinical use for many inflammatory associated diseases. Studies to determine if such an approach may be useful to prevent cognitive dysfunction but not inhibit the cancer chemotherapeutic properties of ADR are in progress.

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