Loss of Caspase-2-dependent Apoptosis Induces Autophagy after Mitochondrial Oxidative Stress in Primary Cultures of Young Adult Cortical Neurons

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Mitochondrial dysfunctions have been associated with neuronal apoptosis and are characteristic of neurodegenerative conditions. Caspases play a central role in apoptosis; however, their involvement in mitochondrial dysfunction-induced neuronal apoptosis remains elusive. In the present report using rotenone, a complex I inhibitor that causes mitochondrial dysfunction, we determined the initiator caspase and its role in cell death in primary cultures of cortical neurons from young adult mice (1–2 months old). By pretreating the cells with a cell-permeable, biotinylated pan-caspase inhibitor that irreversibly binds to and traps the active caspase, we identified caspase-2 as an initiator caspase activated in rotenone-treated primary neurons. Loss of caspase-2 inhibited rotenone-induced apoptosis; however, these neurons underwent a delayed cell death by necrosis. We further found that caspase-2 acts upstream of mitochondria to mediate rotenone-induced apoptosis in neurons. The loss of caspase-2 significantly inhibited rotenone-induced activation of Bid and Bax and the release of cytochrome c and apoptosis inducing factor from mitochondria. Rotenone-induced downstream activation of caspase-3 and caspase-9 were also inhibited in the neurons lacking caspase-2. Autophagy was enhanced in caspase-2 knock-out neurons after rotenone treatment, and this response was important in prolonging neuronal survival. In summary, the present study identifies a novel function of caspase-2 in mitochondrial oxidative stress-induced apoptosis in neurons cultured from young adult mice.

Mitochondrial dysfunction due to mutations or exposure to toxic agents leads to cellular abnormalities, including decreased ATP synthesis and increased generation of reactive oxygen species (ROS) that may result in cell death (1). There is strong evidence suggesting that mitochondrial dysfunction plays an important role in neurodegeneration and neuronal death in conditions like Alzheimer disease, Parkinson disease, amyotrophic lateral sclerosis, and Huntington disease (1–3). As a result of these neurodegenerative insults, a major portion of the affected neurons dies through a process known as apoptosis (4). Apoptosis is the programmed cell death pathway mainly executed by cysteine proteases known as caspases. The apoptotic cascade of caspses is initiated by the activation of apical (initiator) caspases that include caspase-2, caspase-8, caspase-9, and caspase-10 (5–7). In response to noxious stimuli and related cellular stress situations, initiator caspases directly or indirectly activate executioner caspses, which in turn orchestrate apoptotic cell death (8). Although a growing amount of evidence suggests that caspase-mediated apoptosis contributes to neuronal cell death in a variety of neurodegenerative conditions, the identification of the caspases that initiate this response remain elusive (9, 10). Furthermore, a role for autophagy has also been identified in neurons during neurodegenerative disorders. Although autophagy generally prevents neuronal cell death, whether it plays a protective or detrimental role in neurodegenerative disease depends on the environment (11).

To gain insight into the mechanisms of neuronal death, selective neurotoxins have been identified that can cause mitochondrial dysfunction and, thus, create cellular and animal models of neurodegeneration. One of the most widely used neurotoxins is rotenone, which inhibits complex I in the mitochondrial oxidative phosphorylation pathway (12, 13). Complex I inhibition results in a decrease in ATP synthesis and an accumulation of oxidative radicals, causing detrimental oxidative stress and cell death (13–15). The mechanism of mitochondrial oxidative stress-induced cell death is well studied in embryonic neurons or cells of non-nervous tissue origin (16–20). However, it remains to be determined whether and how post-mitotic neurons are affected by mitochondrial oxidative stress-inducing agents. The existing evidence suggests that cell death in post-mitotic neurons involve different pathways than those used by embryonic neurons or the cells of other origin (21). To determine therapeutic approaches for neurodegenerative diseases, it is important to gain a better understanding of the cell death mechanisms induced in well differentiated neurons. Thus, in the present study we cultured neurons from young adult mice and used them as a model to study the mechanism of cell death induced by mitochondrial dysfunction-causing agent, rotenone. Using a technique that traps the initiator caspases in situ, we identified caspase-2 as the initiator caspase that acted upstream of mitochondria to mediate rotenone-induced apoptosis in neurons. Loss of caspase-2 inhibited rotenone-induced apoptosis. Inhibition of apoptosis was associated...
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

associated with increased autophagy in rotenone-treated neurons lacking caspase-2, which was associated with an increased survival. Thus, the present study identifies a novel function of caspase-2 in mitochondrial oxidative stress-induced apoptosis in neurons cultured from young mice.

**EXPERIMENTAL PROCEDURES**

*Reagents—Glutamine, B27 supplement, fibroblast growth factor-2, penicillin/streptomycin, and Neurobasal A media were purchased from Invitrogen. Hibernate A was from BrainBits (LLC). Laminin, OptiPrep 1.32, papain, poly-D-lysine, rotenone, N-acetylcysteine (NAC), and trypan blue were purchased from Sigma. Pepstatin A, 3-methyladenine, bafilomycin A1, and E-64d were from Calbiochem. AlexaFluor 488 or Alexa Fluor 568 IgG secondary antibody, Calcein-AM, Hoehcht 33258, propidium iodide (PI), LysoTracker Red, and MitoTracker Red CMH2xROS were purchased from Molecular Probes Inc. (Eugene, OR). Vectashield mounting medium was from Vector Laboratories, Inc (Burlingame, CA). The BCA protein assay kit and chemiluminescence system were from Pierce. β-Actin antibody, radiolabeled precipitation assay buffer, and HRP-labeled anti-mouse or anti-rabbit IgG secondary antibodies were from Santa Cruz Biotechnology (Santa Cruz, CA). Antibodies against cytochrome c and Beclin1 were from BD PharmingenTM (BD Pharmingen). Anti-Bax antibody (6A7 mouse monoclonal), Bid, and acetyl-CoA carboxylase were from Abcam (Cambridge, MA). Primary antibodies for apoptosis inducing factor (AIF), glyceraldehyde 3-phosphate dehydrogenase, LC3, caspase-9, caspase-3, and active caspase-3 were from Cell Signaling (Cell Signaling Technology, Danvers, MA). Biotin-Val-Ala-Asp(OMe)-fluoromethyl ketone (b-VAD-fmk) was from Enzyme Systems Products (Livermore, CA), and caspase-2 monoclonal antibody (clone 11B4) was from Alexis Biochemicals (San Diego, CA). In case the source is not mentioned, the chemicals were purchased from Sigma. All the cell culture plastic and glassware were from Nunc (Nalgene Nunc International, Rochester, NY).

*Mice—The caspase-2 knock-out (casp2−/−) mice were originally generated by Dr.Junying Yuan of Harvard University and kindly provided by Dr. Carol Troy of Columbia University with the consent of Dr. Yuan (22). The mice were backcrossed with C57Bl/6 for 10 generations. The mice were housed in microisolation-topped cages and maintained in a pathogen-free environment at the Assessment and Accreditation of Laboratory Animal Care-accredited University of Texas Health Science Center-San Antonio animal facility following the NIH Guidelines for the Care and Use of Laboratory Animals. The University Institutional Animal Care and Use Committee approved the protocols used in this study. The wild type (wt) and casp2−/− animals were maintained in an animal room with controlled temperature and exposed to a 12:12 light-dark lighting regimen with the lights on daily from 06:00 to 18:00. Food was available ad libitum.

**Primary Culture of Cortical Neurons**— Cultures of cortical neurons were established from young adult mice that were 1–2 months old following the protocol published by Brewer and Torricelli (23). Briefly, the brains of the mice were dissected to isolate cortices that were transferred to 4 °C in Hibernate A supplemented with B27. Cortices were diced (0.5 mm thick) and then incubated in media containing papain (12 mg/6 ml in Hibernate A) at 30 °C in a shaker for 30 min. The digested tissue was washed free of papain and triturated 12–15 times using a siliconized Pasteur pipette. The triturated tissues were applied on the top of an OptiPrep 1.32 gradient and centrifuged at 800 × g for 10 min. Debris above 4 ml was discarded, and the fractions containing neurons (fractions 2 and 3) were collected and diluted in 5 ml of Hibernate A/B27. After centrifugation at 200 × g for 2 min, the resulting pellets were resuspended in 3 ml of conditional medium (Neurobasal A/B27, 0.5 mM glutamine, 10 ng/ml fibroblast growth factor-2 and penicillin/streptomycin), and viability was determined by trypan blue exclusion. For cell plating, glass coverslips in 24 multi-well plates were coated with 100 μg/ml poly-D-lysine and laminin. Approximately, 1.25 × 10⁴ cells were plated on each 15-mm glass coverslip and allowed to attach and attain morphology. Purity of neuronal cultures was determined by staining with NeuN antibody or MAP antibody (neuron-specific) and GFAP antibody (astrocyte-specific). The cultures stained >92% for neuronal cell types after 4–7 days of culture (data not shown). Drug treatments were initiated after 5 or 6 days in vitro. For the treatments, the B27 supplement was replaced with B27 lacking antioxidants. Rotenone was added directly to the conditional medium and was not washed out until the end of the experiment. The control neurons were treated with similar amounts of the DMSO (rotenone carrier).

**Culture of Mouse Embryonic Fibroblasts (MEFs)—wt, bak−/−, bak−/−, bak−/− bak−/−, and casp2−/− MEFs were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% FBS, 100 units/ml penicillin, and 100 μg/ml streptomycin.**

**Neuronal Viability and Apoptosis Assay**—To determine neuronal survival, neurons were stained with Calcein-AM (2 μM) and PI (25 μg/ml) for 30 min, and the fraction of live (Calcein-AM-positive) and dead (PI) cells was determined. Hoechst 33258 staining (10 μg/ml, was performed for 30 min to visualize the nuclei. Based on nuclear morphology, cells showing apoptotic features (nuclear condensation or nuclear fragmentation) were scored. Neurons were visualized on a laser-scanning confocal microscope (LSM) (FV1000; Olympus, Tokyo, Japan), and images were captured at 40× magnification. Images were scored by an observer blind to treatment, and a minimum of 200 cells was analyzed per group in triplicate.

**Measurements of Mitochondrial Superoxide**—Mitochondrial superoxide levels were determined using MitoSOX Red (500 nM). Cortical neurons were cultured on coverslip-bottom chamber slides and maintained in Neurobasal A media supplemented with B27 for 6 days in vitro. Cultures were then washed with Krebs-Ringer buffer and incubated for 1 h at 37 °C with either vehicle or rotenone (20 min after MitoSOX Red loading). Subsequently, neurons were rinsed three times with warm Krebs-Ringer buffer and immediately visualized on LSM (FV1000; Olympus). Images were acquired using a 563-nm He-Ne laser to excite MitoSOX Red fluorescence. All images were scanned using 60× magnifications and using the same parameters such as laser power, detector gain, offset, pinhole,
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

and zoom to maintain standardized imaging conditions. Analysis of fluorescence data was performed with ImageJ (Windows version; National Institutes of Health). An appropriate threshold was employed to eliminate background signal in the images before histogram analysis.

Immunofluorescence Staining for Cytochrome c, AIF, Bax, Active Caspase-3, and LC3—To visualize mitochondria or lysosomes, live neurons were stained with MitoTracker Red (250 nM) or LysoTracker Red (500 nM) for 30 min and were washed free of dye before fixation. The neurons were fixed by the addition of 3.7% paraformaldehyde for 20 min at room temperature. Cells were washed, permeabilized (0.3% Triton X-100), blocked in 1% BSA for 1 h at room temperature, and immunolabeled with the primary antibody (1:200 dilution and overnight incubation at 4 °C). Neurons were then washed with PBS and incubated for 2 h with AlexaFluor 488 or Alexa Fluor 568 IgG secondary antibody (1:1000 dilution) at room temperature. For nuclear staining, Hoechst 33258 (10 µg/ml) was added along with the secondary antibody. Cells were washed with PBS and then with water. The coverslips were mounted with Vectashield mounting medium and sealed. Approximately, 150–200 neurons per coverslip were visualized on LSM (Olympus FV1000), and images were digitally captured and scored blindly. Each experiment was repeated at least three times with separate batches of neuronal cultures.

In situ Labeling of Active Caspases—To capture the active initiator caspase, neurons were pretreated with cell-permeable b-VAD-fmk (50 µM) for 2 h in the conditional media. Rotenone was then added to the neuronal cultures, and samples were collected at selected time points. To study the total active caspses, the neurons were treated with rotenone for 36 h and were incubated for 2 h with b-VAD-fmk (50 µM) before harvest. The neurons were pelleted and lysed by freeze-thawing (5–6 times) in 500 µl of CHAPS lysis buffer (150 mM KCl, 50 mM HEPES, 0.1% CHAPS, and 0.1% Nonidet P-40 with pH 7.4 also containing the protease inhibitors (2 µg/ml) and phenylmethyisulfonyl fluoride). After centrifugation at 15,000 × g for 10 min, the supernatant was collected, and the b-VAD-fmk-bound active caspases were captured on 30 µl of streptavidin-conjugated agarose beads. After overnight rotation at 4 °C, the agarose beads were extensively washed in lysis buffer. The biotinylated proteins were eluted from the beads by the addition of SDS sample buffer and incubated at 95 °C for 10 min. Whole cell lysates (WCL) (25 µg) that did not go through streptavidin precipitation or the eluted biotinylated proteins (25 µl) were used for SDS-PAGE and subsequent Western blotting. The endogenously biotinylated protein acetyl-CoA carboxylase functioned as a control for both pulldown efficiency and loading.

Western Blotting—Protein samples were separated on 4–12% gradient SDS-PAGE and transferred to PVDF membranes. The membranes were blocked in 5% skin milk for 1 h and probed overnight at 4 °C with primary antibodies in 1% skin milk or BSA. The membranes were washed and incubated for 1 h with appropriate HRP-conjugated secondary antibodies. Blots were washed and developed by using an enhanced chemiluminescence system according to manufacturer’s instructions and exposed to Hyperfilm-ECL (Amersham Biosciences). The same blots were reprobed with β-actin or GAPDH or β-tubulin antibody as a loading control (no change in β-actin or GAPDH or β-tubulin protein levels was seen after treatment). Densitometry analysis band intensity was performed using Image J software.

Data Analysis—Data in graphs are reported as ± S.E. and depict the average of at least three independent experiments. Morphological images are representative of at least three independent experiments with similar results. Statistical analysis was performed by Student’s paired t test or a one-way analysis of variance followed by a Bonferroni post test when appropriate (p < 0.05 was considered significant).

RESULTS

Rotenone Induces Apoptosis in Cultures of Cortical Neurons from Young Adult Mice—The dose-dependent effects of rotenone treatment have not been studied in cultures of cortical neurons obtained from young adult mice. Thus, we first determined the dose-dependent cytotoxic effect of rotenone in neuronal cultures from young adult mice. The neurons were treated with different concentrations of rotenone (0.01–1 µM) for 36 h, and viability was determined by staining neurons with Calcein-AM and PI (Fig. 1A). Rotenone induced a dose-dependent increase in neuronal death, and 0.3 µM rotenone, which resulted in about 48% neuronal death, was used for further experiments. Time course studies showed that rotenone induced changes in neuronal morphology within 4 h of treatment, and the fraction of dead cells began to increase after 16 h (~29%) and continued to rise over the next 72 h (see Fig. 3B). To identify the mode of cell death, the nuclei were stained with Hoechst 33258. We observed that rotenone treatment induced apoptosis in neurons that was characterized by chromatin condensation and/or nuclear fragmentation (Fig. 1, B and C).

Caspase-2 Is the Initiator Caspase Activated during Rotenone-induced Apoptosis—Because caspases are the major executors of apoptosis, we determined which caspase acts as an initiator caspase in rotenone-treated cortical neurons. Caspase-2, caspase-8, and caspase-9 are the known initiator caspses (5–7), and the type of cellular stress determines which caspase acts as the initiator in the apoptotic process (24). To identify the initiator caspase, we employed the in situ trapping approach (6). Using a biotinylated form of the caspase inhibitor (b-VAD-fmk) that binds irreversibly to activated caspase, active caspase can be isolated from cell lysates and other in vitro preparations with the use of streptavidin (25). Incubation of the cells with b-VAD-fmk before rotenone treatment leads to trapping of the active initiator caspase and also prevents the activation of the downstream caspases. Therefore, the caspase bound to the biotinylated caspases inhibitor can be identified as the initiator caspase. We treated neurons with b-VAD-fmk (50 µM) for 2 h before adding rotenone. After rotenone treatment, neurons were harvested at different times; the initiator caspase was precipitated using streptavidin. Western blotting was performed to identify the bound caspase by probing the membranes with known initiator caspase antibodies, i.e., anti-caspase-2, caspase-8, and caspase-9 antibodies; the results are shown in Fig. 2A, and the WCL that did not undergo streptavidin capture are shown in Fig. 2B for comparison. The addition of b-VAD-fmk inhibited the apoptotic nuclear mor-
phology that was induced by rotenone treatment (data not shown). The results demonstrate that caspase-2 was pulled down by streptavidin precipitation within hours of the initiation of treatment, suggesting its function as the initiator caspase. Interestingly, besides observing the 47/49-kDa form that was expected during the early hours of rotenone treatment, the 12/14-kDa-cleaved form of caspase-2 was also identified. However, with more extended time, an increase in the 47/49-kDa form of caspase-2 was observed, indicating its activation (Fig. 2Aa). It was interesting to note that even in the WCL, within early hours of rotenone treatment the proform of caspase-2 was cleaved with the appearance of the different cleaved forms that increased further with time (Fig. 2Ab). We did not detect caspase-8a/b forms (55/57 kDa) or caspase-9 (Fig. 2Ac) in our immunoblots (performed on streptavidin pulldown samples) even after a longer time of exposure that also corroborated with WCL where no cleavage or reduction in the proform of caspase-8 or caspase-9 was observed (Fig. 2B), suggesting that these caspases are not the initiators during rotenone-induced apoptosis in neurons. By contrast, the specific engagement of caspase-8 or caspase-9 was observed in primary activated neurons pretreated with b-VAD-fmk and subjected to anti-Fas treatment or staurosporin, respectively (supplemental Fig. 1). Thus, our results identify caspase-2 as an initiator caspase in rotenone-induced apoptosis in primary cortical neurons.

**Loss of Caspase-2 Inhibits Rotenone-induced Apoptotic Cell Death**—Next, we determined whether caspase-2 contributes to rotenone-induced neuronal apoptosis. To avoid potential problems associated with transfection and knockdown using
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

To determine whether the delayed cell death in the rotenone (0.3 μM) for 36 h, in contrast, similarly treated wt neurons after 36 h, in contrast, similarly treated wt neurons as compared with wt. Although rotenone treatment induced apoptosis in nearly 55% of the wt neurons after 36 h, in contrast, similarly treated casp2−/− neurons did not show a significant increase in apoptosis. To determine whether the casp2−/− neurons were dying via an alternative (nonapoptotic) mechanism, we also measured neuronal survival in casp2−/− neurons by Calcein-AM/PI staining (live/dead assay). Interestingly, after rotenone exposure, casp2−/− neurons showed a survival advantage (p < 0.05) as compared with wt (Fig. 3B). Even after 72 h of rotenone treatment, ~68% of wt neurons died compared with only about ~30% of the casp2−/− neurons. When we studied the mode of the delayed cell death in the casp2−/− neurons, the majority of the neuronal morphologies showed a preserved chromatin structure but with the loss of neurites and membrane damage (loss of Calcein-AM and PI staining without showing nuclear condensation or fragmentation), characteristics suggestive of necrosis (Fig. 3C). Taken together, these results indicate that after treatment with rotenone, caspase-2 is activated as the initiator caspase, and loss of caspase-2 is protective against rotenone-induced apoptosis as well as neuronal death.

FIGURE 3. Caspase-2 is essential for rotenone-induced neuronal apoptosis. A, cortical neurons cultured from wt and casp2−/− mice were treated with rotenone (0.3 μM) for 36 h. Apoptotic cells were determined on the basis of nuclear morphology visualized by Hoechst 33258 staining. B, cell viability was determined by staining live cells with Calcein-AM and dead cells with PI. The time course of rotenone treatment (0.3 μM) shows cell death % control in wt and casp2−/− cortical neurons at different time points. Data (mean ± S.E., n = 150–200 neurons) were derived from three independent experiments with different batches of cortical neurons. Asterisks indicate a significant change (protection) comparing wt versus casp2−/− cortical neurons (*, p < 0.05). C, representative images of neurons treated with rotenone for 0, 36, or 72 h and stained with Hoechst 33258 (blue, indicating total nuclei), Calcein-AM (green, indicating live cells), and PI (red, indicating dead cells). Images (LSM) were taken with a 40× objective, repeated three times with different cultures.

Loss of Caspase-2 Protects Neurons from Rotenone-induced Superoxide—To determine whether rotenone-induced mitochondrial superoxide generation is responsible for caspase-2 activation and cytotoxicity, we conducted studies using a well known antioxidant, NAC, that is known to inhibit rotenone-induced superoxide (18, 19, 26). We found that rotenone induced superoxide generation, as determined by MitoSOX Red staining, in neuronal cultures within 1 h of treatment. No significant difference was observed in the extent of superoxide production in rotenone-treated wt and casp2−/− after rotenone treatment. Pretreatment with NAC significantly inhibited rotenone-induced superoxide generation (Fig. 4A) as well as cell death in the wt neurons (Fig. 4B), confirming a role for rotenone-induced superoxide generation in inducing neuronal cell death.

We further investigated the effect of NAC pretreatment on rotenone-induced caspase-2 cleavage (Fig. 4C). As determined by Western blotting, NAC treatment significantly inhibited the rotenone-induced caspase-2 cleavage. We then studied the effect of NAC treatment on rotenone-induced cell death in casp2−/− neuronal cultures (Fig. 4C). Interestingly, pretreatment with NAC did lead to significant inhibition of superoxide production in rotenone-treated casp2−/− neurons but did not provide any additional protection to casp2−/− neurons from rotenone-induced cell death (Fig. 4D). Collectively, the data suggest that caspase-2 is activated by rotenone-induced superoxide and plays a role in neuronal cell death.
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

We further investigated whether rotenone stimulates superoxide production and caspase-2 activation in another model and, if so, whether caspase-2 is required for cell death. For this we performed experiments on MEF from wt or casp2−/− mice (supplemental Fig. 2). Despite a similar extent of rotenone-induced superoxide production as determined by MitoSOX staining, casp2−/− MEFs showed significantly higher survival against rotenone treatment, suggesting that caspase-2 plays a role in superoxide-mediated cell death not only in neurons but also in other cell types.

Caspase-2 Acts Upstream of Rotenone-induced Mitochondrial Apoptotic Events—We further investigated how caspase-2 is involved in rotenone-induced apoptosis. Caspase-2 has been shown to induce mitochondrial outer membrane permeabilization (MOMP) that leads to the release of pro-apoptotic molecules from mitochondria (27–30). It is well documented that rotenone-induced mitochondria, cytochrome c staining was still punctate and retained its mitochondrial localization (Fig. 5Ad). Quantitative analysis of the percentage of cells showing cytochrome c release confirmed that rotenone-induced cytochrome c release was significantly reduced in casp2−/− neurons compared with wt (Fig. 5B). In line with these data, we also found that loss of caspase-2 protected neurons from rotenone-induced loss of mitochondrial membrane permeability transition as determined by tetramethylrhodamine methyl ester staining (supplemental Fig. 3).

Next, we examined AIF nuclear translocation in rotenone-treated wt and casp2−/− neurons. AIF is another proapoptotic molecule that is released from the mitochondrial intermembranous space after MOMP. After its release, AIF translocates to the nucleus, where it induces large scale DNA fragmentation and chromatin condensation (32–34). To study AIF nuclear translocation, we performed immunofluorescence staining using AIF antibody (red); nuclei were stained with Hoechst 33258 (blue) (Fig. 5C). The quantification of the percentage of cells showing AIF nuclear translocation is shown in Fig. 5D. In the control wt or casp2−/− neurons, AIF retained its mitochondrial localization, as can be seen by its punctate staining pattern (Fig. 5C, a and e). After 16 h of rotenone treatment, cells did not exhibit any significant release of AIF into the cytosol or the nucleus in either wt or casp2−/− neurons (Fig. 5C, b and f). After 24 h of rotenone treatment, there was a significant increase in the percentage of wt neurons (~22%) in which AIF showed loss of punctate staining pattern with a concomitant increase in nuclear and cytoplasmic localization. This pattern in wt increased further after 36 h when more than 51% neurons
showed AIF nuclear translocation (Fig. 5Cd). In contrast, even after 36 h of rotenone treatment only a few nuclei in the casp2−/− neurons showed nuclear localization of AIF (~11%). This response was significantly lower than that observed in wt (Fig. 5C, d and h), suggesting an involvement of caspase-2 upstream to AIF release in rotenone-induced neuronal apoptosis. Thus, the results indicate a role of caspase-2 in the induction of MOMP during rotenone-induced toxicity in neurons.

One way by which caspase-2 can induce MOMP is by activating Bid and Bax (28, 35). We examined whether rotenone treatment leads to Bid cleavage and, if so, whether this response is affected by loss of caspase-2. In wt neurons, rotenone treatment induced the truncation of Bid with a decrease in the 21-kDa form and a subsequent increase in the 15-kDa form, as determined by Western blotting (Fig. 6A). However, compared with wt, the truncation of Bid was significantly reduced in rotenone-treated casp2−/− neurons, suggesting an involvement of caspase-2 in rotenone-induced Bid truncation.

We next examined rotenone-induced Bax activation in wt and casp2−/− neurons by immunolabeling the cells using the Bax 6A7 antibody that recognizes the active Bax (Fig. 6B). In both wt and casp2−/− controls, no significant Bax 6A7 fluorescence staining was detected, indicating that in untreated neurons Bax was not present in the active form (Fig. 6B, a and c). After rotenone treatment, there was a significant increase in the number of wt neurons exhibiting Bax 6A7 staining compared with the controls (Fig. 6Bh), suggesting Bax activation. In contrast, very few rotenone-treated casp2−/− neurons exhibited Bax 6A7 staining (Fig. 6Bd). Quantitative analysis of Bax-stained cells showed that rotenone-induced Bax activation was significantly less in casp2−/− neurons compared with that in wt (Fig. 6C).

**FIGURE 5. Loss of caspase-2 prevents rotenone-induced release of cytochrome c and AIF from mitochondria.** Cortical neurons cultured from wt and casp2−/− were treated with rotenone (0.3 μM) for 16 h. A, neurons were co-stained with MitoTracker Red for mitochondria, cytochrome c (green), and Hoechst 33258 for nuclei (blue), and overlay images are shown. Mitochondrial localization of cytochrome c can be seen as yellow fluorescence due to colocalization of green fluorescence with red (mitochondria). Cytochrome c release can be identified as an increase in green fluorescence with a diffused pattern and a decrease in the yellow fluorescence localized to mitochondria. Images were obtained with a 60× objective, shown as 3 times the digital magnification. B, a histogram shows the % of cells showing cytochrome c translocation to cytosol. Loss of caspase-2 prevents rotenone-induced nuclear translocation of AIF. C, cortical neurons cultured from wt and casp2−/− were treated with rotenone (0.3 μM) for 16, 24, and 36 h and were stained for AIF (red) and nuclei-stained with Hoechst 33258 (blue). The overlay images of AIF and nuclear staining are shown at 60× magnification (2× digital zoom). The AIF translocation to the nuclei can be identified by the appearance of a magenta color due to co-localization with blue fluorescence (nuclei) with red (AIF). D, a histogram shows the % of cells showing AIF nuclear translocation. At least 150–200 neurons were counted per group. Data (± S.E.) were derived from three separate experiments. The asterisk (*) represents p values <0.05, comparing time-matched wt versus casp2−/− neurons.
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

Caspase-2 Activation Is Upstream of Caspase-9 and Caspase-3 Activation—Release of cytochrome c from mitochondria results in formation of the apoptosome with a subsequent activation of caspase-9 and caspase-3. Because our earlier results indicated that caspase-2 is an initiator caspase during rotenone-induced apoptosis, we next examined rotenone-induced activation of caspase-9 and caspase-3 in wt and casp2−/− neurons. To detect only active caspases, after 36 h of rotenone treatment, the WCL were incubated with b-VAD-fmk. The caspases trapped by the inhibitor were isolated using streptavidin pulldown and then analyzed by Western blotting. The WCL without streptavidin pulldown are also shown for comparison (Fig. 7B). After treatment with rotenone for 36 h, there was a significant increase in the activation of caspase-9 and -3 in the wt neurons (Fig. 7A). By contrast, in the casp2−/− neurons, the levels of active caspase-3 and -9 were detected only at very low levels. We could not detect the activation of caspase-8 either in wt or casp2−/− neurons after rotenone treatment (data not shown). We also performed immunofluorescence microscopy using an antibody that detects cleaved caspase-3 (active subunit) followed by its quantification (Fig. 7C and D). In untreated wt neurons, very few cells showed active caspase-3 (∼6% in wt and ∼7% in casp2−/−). After treatment with rotenone for 36 h, ∼47% of the wt neurons stained positive for the cleaved caspase-3 (Fig. 7Cb). By contrast, no significant increase in active caspase 3 was observed in the rotenone-treated casp2−/− neurons (Fig. 7Cd).

The use of a commercial caspase inhibitor showed that inhibition of caspase-2 provided maximum protection from rotenone-induced neuronal death. On the other hand, treatment with caspase-9 or caspase-3 inhibitors produced less protection of the neurons from rotenone toxicity. Similarly, the caspase-8 inhibitor did not provide any significant protection from neuronal cell death (supplemental Fig. 4). Collectively, these results suggest a selective initiator role for caspase-2 in rotenone-induced neuronal apoptosis.

Autophagy Is Induced in casp2−/− Neurons and Delays Rotenone-induced Cell Death—Loss of caspase-2 inhibited rotenone-induced apoptosis and delayed cell death in the cortical neurons. Therefore, it was important to determine whether casp2−/− neurons survive better in the presence of rotenone-induced oxidative damage or if the remaining casp2−/− neurons show a delayed cell death by an alternative pathway in the absence of apoptosis. We and others have shown that, besides inducing apoptosis, oxidative stress can also induce autophagy (26, 36–40). Thus, we examined whether rotenone induces autophagy in casp2−/− neurons. Autophagy can be detected by examining the conversion of cytosolic LC3-I to lipidated, autophagosome membrane-bound LC3-II. These autophagosomes in turn form autophagolysosomes by fusing with lysosomes (41). Immunocytochemistry using LC3 antibody was performed to study its redistribution after rotenone treatment. In addition, LysoTracker Red colabeling was performed to visualize fusion between lysosomes and autophagosomes (Fig. 8A).

In the control wt and casp2−/− neurons, LC3 was distributed predominantly as diffuse green fluorescence in the cytoplasm and nucleus with very little lysosomal staining. After rotenone treatment, wt neurons retained a diffused staining pattern of

FIGURE 6. Loss of caspase-2 prevents rotenone-induced Bid cleavage and Bax activation. Cortical neurons cultured from wt and casp2−/− were treated with rotenone (0.3 μM) for the indicated times. A, a representative Western blot performed using WCL shows Bid cleavage in rotenone-treated wt neurons (36 h treatment) that was prevented in the absence of caspase-2. Full-length Bid appears at 21 kDa, and cleaved Bid is at 15 kDa. B, immunocytochemistry for Bax translocation using Bax 6A7 (green) antibody that recognizes active Bax is shown. Overlay images of Bax staining and nuclei (Hoechst 33258 in blue) are shown (60×). Bax staining was observed only when active Bax was present. C, a histogram shows the % of cells showing Bax activation (±S.E., n = 150–200 neurons/group); the experiment was repeated three times. The asterisk (*) represents p values <0.05, comparing time matched wt versus casp2−/− neurons.
LC3 with very little (non significant) increase in lysosomal staining (Fig. 8A). By contrast, rotenone-treated casp2−/− neurons at 16 h showed a characteristic punctate pattern of LC3 immunofluorescence that was dispersed through the cell. Furthermore, a fraction of the casp2−/− neurons demonstrated a colocalization of LC3 with lysosomes, suggesting the formation of autophagosome and autophagolysosomes. After 36 h of rotenone treatment of casp2−/− neurons, the punctate feature representing the autophagosomes gradually faded/vanished, presumably due to the fusion of autophagosome with lysosome resulting in degradation of LC3 and an associated loss of staining.

We also performed Western blotting for LC3. Because LC3-II is continually degraded within autophagolysosomes during autophagy, we treated the neurons in the presence of pepstatin A (1 μM) and E-64d (10 μM), which are lysosomal inhibitor, and the samples were prepared for Western blotting for LC3 to demonstrate autophagic flux (42). A significant increase in LC3-II was detected in rotenone-treated casp2−/− neurons but not in wt neurons, further corroborating the evidence of enhanced autophagy in the rotenone-treated casp2−/− neurons (Fig. 8, B and C). We also performed Western blotting for Beclin1, a component of the phosphatidylinositol 3-kinase complex known to increase during autophagy (43). A significant and consistent increase was observed in the levels of Beclin1 in rotenone-treated casp2−/− neurons as compared with wt neurons after rotenone treatment. It is important to note that in rotenone-treated wt neurons, Beclin1 levels decreased significantly (Fig. 8, D and E). It has been shown that Beclin1 is cleaved by caspases during apoptosis (44), which is consistent with our data that indicate an induction of apoptosis in wt neurons after treatment with rotenone.

Next, we examined the role of autophagy on cell death or cell survival. Because transfecting the young adult neurons in cul-

**FIGURE 7.** Effect of rotenone on caspase-9 and caspase-3 activation in wt and casp2−/− neurons. Neurons from wt and casp2−/− cultures were either untreated or exposed to rotenone. A and B, neurons were incubated with rotenone for ~36 h, and b-VAD-fmk (50 μM) was added 2 h before harvesting. Cell lysates were subjected to streptavidin pulldown (A) and/or preparation of WCL (B) followed by assessment for caspase-2, -3, and -9 by immunoblotting. Western blot analysis was performed by standard procedures. The same blots were subsequently probed with acetyl-CoA carboxylase (ACOC) or β-actin and used as a control for precipitation and loading. Shown are representative blots; the experiment was repeated three times. C, shown is immunofluorescence staining for active caspase-3 (green) and nuclei counterstained with Hoechst 33258 (blue). Shown are the representative overlay images at 60× magnifications. D, a histogram shows the % of cells with active caspase-3 staining, quantified by counting the numbers of cells with active caspase-3 and the total number of cells (±S.E., n = 150–200 neurons/group); the experiment was repeated three times. The asterisk (*) represents p values <0.05, comparing time matched wt versus casp2−/− neurons.
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

ture was associated with potential technical difficulties, we were restricted to the use of pharmacological inhibitors of autophagy. The inhibition of autophagy at an early stage by 3-methyladenine and at a late stage by bafilomycin A1 completely abolished the protective effect of loss of caspase-2 during rotenone-induced cell death (Fig. 8F). The enhanced death

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**Figure 8**

A. Images showing the effects of rotenone on LC-3 staining in wild-type (wt) and caspase-2 knockout (casp2−/−) neurons. (a) Control wt; (b) 16 hr of rotenone; (c) 36 hr of rotenone; (d) control casp2−/−; (e) 16 hr of rotenone; (f) 36 hr of rotenone. Scale bar: 10 μm.

B. Western blot analysis of LC3-I and LC3-II in control and rotenone-treated wt and casp2−/− neurons.

D. Western blot analysis of Beclin 1 in control and rotenone-treated wt and casp2−/− neurons.

G. Western blot analysis of LC3-I and LC3-II in wt and casp2−/− neurons with or without NAC.

H. Western blot analysis of ratio of LC3-I/LC3-II in wt and casp2−/− neurons with or without NAC.

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**Figure 8**

C. Graph showing the ratio of LC3-II/LC3-I in control and rotenone-treated wt and casp2−/− neurons.

E. Graph showing the fold change in Beclin protein intensity compared to control in wt and casp2−/− neurons.

F. Graph showing the cell death % control in wt and casp2−/− neurons with control, rotenone, 3-MA, Baf-A1, rotenone+3-MA, and rotenone+Baf-A1 treatment.

H. Graph showing the ratio of LC3-I/LC3-II in wt and casp2−/− neurons with or without NAC.

J. Graph showing the fold change in Beclin protein intensity compared to control in wt and casp2−/− neurons with or without NAC.
in the absence of caspase-2 was necrotic in nature as determined by an increase in PI uptake with loss of Calcein-AM and without features of apoptosis, such as chromatin condensation and nuclear fragmentation. Thus, loss of caspase-2 results in enhanced autophagy after rotenone treatment and thereby delays neuronal death.

Finally, to find whether rotenone-mediated oxidative stress causes enhanced autophagy in casp2−/− neurons, we accessed autophagy in the presence of NAC (an antioxidant) in rotenone-treated casp2−/− neurons. Autophagy was determined by Western blotting to measure the LC3-II/LC3-I ratio and levels of Beclin1. As shown in Fig. 8, G–J, we found that NAC pretreatment inhibited rotenone-induced increase in LC3-II/LC3-I ratio as well as Beclin1 levels in casp2−/− neurons, indicating that NAC inhibits rotenone-induced autophagy. These results suggest that autophagy was induced in response to oxidative stress induced by rotenone when the neurons cannot undergo apoptosis due to lack of caspase-2 (Fig. 9).

**DISCUSSION**

The function of caspase-2 has remained enigmatic, partly due to the fact that mice lacking caspase-2 are viable and fertile due to the fact that mice lacking caspase-2 are viable and fertile and without features of apoptosis, such as chromatin condensation and nuclear fragmentation. Thus, loss of caspase-2 results in enhanced autophagy after rotenone treatment and thereby delays neuronal death.

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**DISCUSSION**

The function of caspase-2 has remained enigmatic, partly due to the fact that mice lacking caspase-2 are viable and fertile with only minor apoptotic defects in certain cell types (22). However, several studies have implicated caspase-2 as a crucial mediator of apoptosis in a context-dependent manner, which includes heat shock (25), DNA damage (29), and cytoskeleton disruption (45). Previous studies from our laboratory demonstrated that casp2−/− mice display premature aging-related traits and significantly higher levels of oxidized proteins in liver than wt mice, indicating an inability to remove the oxidatively damaged cells (46). Furthermore, caspase-2 functions in oxidative stress-induced apoptosis are coming to light (47, 48). In this regard a present report provides strong evidence for the role of caspase-2 in mitochondrial-mediated oxidative stress-induced apoptosis in neurons. Moreover, involvement of caspase-2 in rotenone-induced cell death is not restrictive to neuronal cell type but also non-neuronal cell type (MEFs).

The present study was performed employing cultures of young adult cortical neurons to identify the initiator caspase and its role in neuronal apoptosis induced by mitochondrial oxidative stress inducing agent, rotenone. Our results demonstrate 1) a role of caspase-2 as an initiator caspase during mitochondrial oxidative stress-induced apoptosis, 2) that caspase-2 acts upstream of mitochondria during rotenone-induced apoptosis, 3) that loss of caspase-2 protect neurons from apoptosis as well as cell death, and 4) that rotenone induces autophagy in casp2−/− neurons that delays the cell death and neurons eventually die via necrotic cell death.

Caspase-2 is one of the most conserved caspases and has been recognized as both an initiator and an effector caspase depending upon the cell type and type of stressor (49–55). In the case of neurons, the initiator role of caspase-2 has remained controversial. Previous studies have defined a role of caspase-2 as an initiator caspase in neuronal apoptosis during serum deprivation (56), β-amyloid-mediated toxicity (57), and oxidative stress-induced apoptosis of neuronal stem cells (47), where it plays an important role in apoptosis induction. On the other hand, other studies have suggested that caspase-2 may be activated in response to different stimuli in neurons and other cell types but may not be essential for induction of apoptosis (22,54). In our experiments employing *in situ* trapping of initiator caspases approach, which is a valuable method to identify initiator caspases (6, 25), we identified caspase-2 as the initiator caspase during rotenone-induced apoptosis in neurons. Furthermore, using neurons that lack caspase-2, our data clearly indicate that caspase-2 plays an important role in mediating neuronal apoptosis during mitochondrial oxidative stress. The initiator role of caspase-2 was further supported by the observations that loss of caspase-2 inhibited rotenone-induced caspase-3 and caspase-9 activation and a caspase-2 inhibitor...
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

provided maximum protection from cell death compared with other caspase inhibitors used in the study.

In the present study we found that rotenone-induced apoptosis in the cultures of young adult cortical neurons was mainly due to its capacity to generate mitochondrial superoxide, which is in agreement with previous observations (18, 19, 26). Interestingly, we also found that caspase-2 activation was rotenone-induced superoxide-dependent. We suggested that if caspase-2 is involved in mitochondrial oxidative stress-induced cell death, then inhibition of superoxide should not provide further protection from cell death in casp2−/− neurons. In line with this hypothesis, pretreatment of casp2−/− neurons with antioxidant did not provide further protection to casp2−/− neurons from rotenone-induced cell death. Thus, our data suggest that caspase-2 is activated by rotenone-induced superoxide and plays a role in apoptotic cell death. However, in the present study we did not determine, 1) how superoxide activates caspase-2 and 2) if there is a particular species of superoxide or ROS that is responsible for caspase-2 activation and needs further investigation. Based on the literature, the structure of caspase-2 carries the maximum number of cysteines among the caspase family that includes a cysteine at the processing site and a central disulfide bridge that leads to dimer stabilization in caspase-2 (58). Cysteine is an oxidative target because of the reactivity of the thiol group that is susceptible to modification by free radicals that may modulate the activity of these proteins, thus making caspase-2 a target for oxidation-based regulation. Furthermore, Madesh et al. showed that incubation with superoxide induced caspase-2 cleavage and, hence, suggested that O2− triggers caspase-2 cleavage through an as yet unidentified mechanism. They also showed that hydrogen peroxide did not recapitulate the effects of superoxide in promoting caspase-2 activation. Moreover, in agreement with these observations, we found that loss of caspase-2 protected against rotenone-induced neuronal cell death but not hydrogen peroxide-induced cell death (data not shown). Together these results indicate that caspase-2 may be more specifically involved in superoxide but not peroxide-mediated cell death and needs further investigation.

Previous studies performed to understand the mechanism of rotenone-induced apoptosis in various models showed an involvement of the intrinsic pathway of apoptosis that leads to activation of caspase-3 and caspase-9 after cytochrome c release (16–20). In agreement with previous observations, our results also demonstrate that neuronal cultures from young adult mice engage the intrinsic pathway of apoptosis. In addition, the present study demonstrates caspase-2 activation as an upstream event that engages the mitochondrial-dependent apoptotic pathway by inducing the release of cytochrome c and AIF from the mitochondria. One way by which caspase-2 has been shown to modulate mitochondrial release of apoptogenic proteins is by modulating the proapoptotic members of Bcl-2 protein family namely, Bid and Bax (28, 35). In the present study we did find inhibition of rotenone-induced Bid and Bax activation in cells lacking caspase-2, further suggesting that caspase-2 induces MOMP in neurons after rotenone treatment. However, previously it has been shown that superoxide-induced MOMP requires Bak for cytochrome c release in a Bak-independent manner. Using the similar system (bax−/−, bak−/−, and bax−/− bak−/− MEFs), indeed we found that despite the difference in the system used to generate superoxide, cytochrome c release was dependent on Bak but not Bax (supplemental Fig. 5).

It has been reported that elimination of certain caspases can be compensated by the activation of other caspases (25, 59, 60). For example, Troy et al. (61) found that caspase-2 plays a predominant role during nerve growth factor deprivation-induced cell death; however, in the absence of caspase-2, a compensatory switch to the caspase-9 pathway is observed (54). In the present study, we found that in the casp2−/− neurons the basal level of caspase-9 was higher than the wt (data not shown), which is consistent with the report by Troy et al. (54). However, we did not observe a compensatory activation of caspase-9 or caspase-8 in response to rotenone treatment in casp2−/− neurons. Moreover, these neurons die a delayed necrotic cell death that further suggested a role of caspase-independent apoptosis. Thus, compensatory caspase activation may depend upon the type of stimulus and the cell type.

Although casp2−/− neurons did not show compensatory caspase activation after rotenone treatment in the present case, they did show induction of autophagy in response to rotenone-mediated oxidative stress. Autophagy, which is a tightly regulated process, leads to degradation of intracellular components via the lysosome (62). It has been shown to be both a cell survival and a cell death pathway (63). Our results define a survival role for autophagy, as inhibition of autophagy abolished the survival advantage of rotenone-treated casp2−/− neurons. On the other hand, we also found that these neurons eventually die a delayed cell death with features of necrosis. It seems that rotenone-treated casp2−/− neurons that cannot undergo apoptotic cell death induce autophagy as an alternate mechanism to combat oxidative stress induced by rotenone for the removal of the damaged organelles and macromolecules. However, eventually autophagic process is overwhelmed by the massive amount of damaged material it is trying to clear. This can lead to cell death through a non-apoptotic mechanism, accounting for a delayed necrotic cell death. Therefore, our data support the hypothesis that autophagy can be both a cell survival mechanism and a cell death pathway depending on the extent of cellular damage and the efficiency of the autophagic process (64). It is important to note that in the present study we did not employ general caspase inhibitors that may produce nonspecific affects that include autophagy activation (65). Our data are based on the inhibition of apoptosis through inhibition of caspase-2 that is involved in the initiation pathway of apoptosis. Our data are in accordance with a previous study in which it was found that the MEFs from double-knock-out bax−/− bak−/− mice fail to undergo apoptosis when treated with death stimulus but instead manifest a massive autophagy followed by delayed cell death (64). Bax and Bak are two multidomain protein members of the Bcl2 family that are required for MOMP, which is one of the decisive steps in apoptotic cell death (66). However, autophagy was not induced when apoptosis was inhibited by other perturbations, such as in apaf1−/− and caspase-9−/− mouse embryonic fibroblasts or the addition of the caspase inhibitor benzylxoyacarbonyl-VAD-fmk to wt mouse embryonic fibroblasts. Thus, along with these observa-
Caspase-2 in Neuronal Mitochondrial Stress-induced Apoptosis

In the presence of necroptosis, our present data support the hypothesis that inhibition of caspase-2 upstream to MOMP may be a decisive factor for the cells to undergo autophagy. Taken together the present study identifies a role of caspase-2 as an initiator caspase in mitochondrial oxidative stress-induced apoptosis in the cultures of young adult cortical neurons. When apoptosis is inhibited at an early stage, the neuron undergoes autophagy that delays cell death (Fig. 9). In cells such as neurons, who have very limited capacity to regenerate, the prevention of any kind of damage or loss is of utmost importance. Thus, understanding the mechanism of cell death and the alternate pathways becomes important in defining the pharmacological approaches to prevent neuronal damage.

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