Acetaldehyde Induces Neurotoxicity *In Vitro* via Oxidative Stress- and Ca^{2+} Imbalance-Mediated Endoplasmic Reticulum Stress

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1. Introduction

Alcohol abuse and alcoholism have become severe social problems. Excessive alcohol consumption can not only lead to extensive tissue damage in organs such as the brain, heart, and liver but also promote disease states, including diabetes mellitus and neuropsychiatric disorders such as Alzheimer’s disease [1]. The brain is one of the major targets of alcohol actions. Alcohol-induced brain damage can produce some potential effects of alcoholism, including cognitive disorders such as learning and memory impairment [2].

The major alcohol metabolites are acetaldehyde (ADE) and acetate, and ADE plays a key role in alcohol-induced neurotoxicity [3]. The conventional view believes that the oxidation of alcohol to ADE can occur in the brain through pathways that involve enzymes including alcohol dehydrogenases (ADHs), catalase, and cytochrome P4502E1 [4]. ADE is catalyzed by acetaldehyde dehydrogenase (ALDH) and is metabolized into acetate with the help of P4502E1. The final metabolites of acetate are carbon dioxide and water [5]. For ALDH, there are three isozymes including ALDH1, ALDH2, and ALDH3. The conversion of ADE was mainly performed by ALDH2 in the mitochondria of hepatocytes [3]. It has been found that the levels of ADE in the blood of individuals with defective ALDH2 are much higher in comparison with those of normal individuals after alcohol ingestion [6]. Unfortunately, 30% to 40% of Asians are deficient in ALDH, resulting in a large accumulation of ADE in the body which cannot be metabolized, so compared to alcohol, ADE may cause greater damage to the human body [7]. Some recent
evidences prove that some of the in vivo and in vitro effects of alcohol are mediated by ADE, which may be a crucial factor in the central effects of alcohol [8–10]. Although there are some advancements in the understanding of ADE’s central effects in recent years [4, 11], the exact mechanism underling ADE’s effects remains largely unknown.

ADE has been proved contributing to many adverse effects of alcohol [12]. Significant behavioral evidences implicate that ADE is involved in the psychopharmacological effects of alcohol [13–15]. At low concentrations, ADE induces euphoria [16] and has been involved in alcohol addiction [17, 18]. In fact, people can be exposed to ADE in their daily lives via different routes, including alcoholic beverages, tobacco smoke, and food additives and fragrances [19]. ADE can damage protein and lipid functions, resulting in DNA damage and mutation [20] and increase of reactive oxygen species (ROS) [21], leading to extensive impairments in DNA damage and mutation [20] and increase of reactive oxygen species (ROS) [21], leading to extensive impairments in DNA damage and mutation [20] and increase of reactive oxygen species (ROS) [21], leading to extensive impairments in DNA damage and mutation [20] and increase of reactive oxygen species (ROS) [21], leading to extensive impairments in DNA damage and mutation [20] and increase of reactive oxygen species (ROS) [21], leading to extensive impairments. Although numerous researches have focused on the role of ADE in different aspects of alcohol effects, there is little study that relates ADE to neurotoxicity. In particular, the action mechanism of ADE is still not fully understood.

Endoplasmic reticulum stress (ERS) resulting from accumulation of unfolded and misfolded proteins within endoplasmic reticulum (ER) lumen has been reported to account for the neuronal death induced by cerebral ischemia, hypoxia, and trauma [24, 25]. ERS and unfolded protein response (UPR) are mediated by three transmembrane ER signaling proteins: pancreatic endoplasmic reticulum kinase (PERK), inositol-requiring enzyme 1 (IRE1), and activating transcription factor 6 (ATF6) [26]. Further, ER is closely related to the redox reaction of the energy center mitochondria, leading to an inseparable relationship between ERS and oxidative stress [27]. In addition, studies suggest that ERS may play an important role in alcohol-induced brain damage [28, 29]. Various stimuli disrupting the homeostasis of cells can cause abnormal protein loading and modification of ER [26]. However, the role of ERS in ADE-induced neurotoxicity remains unclear.

In this study, we investigated the cytotoxic effect of ADE in mouse hippocampal HT22 cells and primary cultured cortical neuronal cells and explored the relationship between ERS and ADE-induced neurotoxicity. We found that ADE induced apoptosis of HT22 cells by upregulating the expression of cleaved caspase-3, cleaved caspase-9, and cleaved caspase-12. ADE treatment led to significant activation of ERS and ROS. In addition, inhibition of ROS with antioxidant N-acetyl-L-cysteine (NAC) and mitigation of ERS with ERS inhibitor sodium phenylbutyrate (4-PBA) obviously suppressed ADE-induced apoptosis and the expression of ERS-related proteins.

2. Experimental Materials and Methods

2.1. Materials and Reagents. ADE with 40% purity was obtained from BBI Life Sciences (Shanghai, China). Dulbecco’s modified Eagle’s medium (DMEM)/F12, B27 supplement, fetal bovine serum (FBS), and DMEM were obtained from Gibco (USA). Annexin V-FITC/PI double-staining and TUNEL Apo-Green detection kits were purchased from BioTools, Inc. (Houston, TX, USA). MTT, acridine orange/ethidium bromide (AO/EB), and 4,6-diamino-2-phenylindol dihydrochloride (DAPI) were from Sigma (USA). Ca2+ kit, 2’,7’-dichlorodihydrofluorescein diacetate (DCFH-DA), and NAC were from Beyotime (Nanjing, China). 4-PBA was purchased from Selleckchem (Houston, TX, USA). Bovine serum albumin (BSA) was from Biosharp (Hefei, China). Rabbit anti-CHOP, anti-GRP78, anti-ATF6, anti-PERK, anti-p-PERK, anti-IRE1, and anti-p-IRE1 IgG were purchased from Bioss (Beijing, China). Rabbit anti-cleaved caspase-9 and anti-cleaved caspase-12 IgG were from Proteintech (Wuhan, China) and Santa Cruz Biotechnology (USA), respectively. Rabbit anti-Bax, anti-Bcl-2, horseradish peroxidase-conjugated, and FITC-conjugated secondary antibodies were from Santa Cruz Biotechnology (USA). Rabbit anti-cleaved caspase-3 was obtained from Cell Signaling Technology (USA). βIII-tubulin was from Sigma (USA). TRIZol reagent Qiagen was purchased from Qiagen (Germany). HiFiScript cDNA synthesis kit and GreenStar qPCR master mix were obtained from Cwbio (China) and Bioneer (South Korea), respectively.

2.2. HT22 Cell Cultures. Mouse hippocampal neuron HT22 cell line was obtained from JENNIO Biological Technology (Guangzhou, China). HT22 cells were cultured in DMEM complemented with 10% FBS, 100 mg/ml streptomycin, and 100 U/ml penicillin and were maintained in a humidified incubator at 37°C and 5% CO2. For each experiment, cells were plated until 70–80% confluence and digested with trypsin.

2.3. Primary Culture of Rat Cortical Neuronal Cells. Primary cultured neurons were prepared as previously described [30]. Neuronal cultures were prepared from the cerebral cortex of neonatal Sprague-Dawley rats of postnatal day 1. Animal procedures were designed to minimize the number of animals required, and appropriate steps were taken to avoid unjustified animal procedures and discomfort. After rats were terminated, their brains were quickly removed and cerebral cortices were dissected in ice-cold DMEM/F12 solution. Subsequently, neurons were dissociated and suspended in complete medium containing DMEM/F12, 15% FBS, 100 U/ml penicillin, and 100 U/ml streptomycin sulfates and counted in a hemocytometer. Approximately 105 cells/ml per well were plated on poly-D-lysine-coated 48-well plates in medium according to different experimental requirements and cultured in the incubator with 5% CO2 at 37°C. Neuronal cell culture medium was changed with serum-free DMEM/F12+2% B27 supplement at 48 h; subsequently, half of the cell culture medium was replaced every 48 h. On the 4th day, cultures were fed with medium containing Ara-C (2.5 mg/ml) for 2 days to prevent growth of glial cells. The neurons were cultured 6 days for experimental use.

2.4. MTT Colorimetric Assay. Cell viability was determined by routine MTT assays [30]. Cell growth inhibition percentage was calculated by SPSS 22.0 software.
2.5. Cell Morphology Changes. 2 × 10^5 cells were seeded per well in a 6-well plate and incubated with ADE for 24 h, and the cells were photographed by an optical microscope (Olympus, Tokyo, Japan). Morphological changes of nuclei after staining with Hoechst 33258 solution for 20 min were observed by fluorescence microscopy (Olympus, Tokyo, Japan).

2.6. Detection of Apoptotic Cells. 2 × 10^5 cells were seeded per well in a 6-well plate and incubated with ADE for 24 h, then the cells were stained with AO/EB mixed solution 20 μl (AO : EB = 1 : 1). The morphological changes of the cells were observed with a fluorescence microscope (Olympus, Tokyo, Japan).

2.7. Apoptosis Analysis. HT22 cells were incubated with various doses of ADE for 24 h, then the cells were washed three times with PBS and stained with Annexin V-FITC and PI at 20–25°C for 20 min. Then, the apoptosis rates were measured by FACSscan flow cytometry (Becton Dickinson, Franklin Lakes, NJ, USA).

2.8. Intracellular Ca^{2+} Measurement. In HT22 cells, fluo-4/aacetoxymethyl ester (AM) can be deacetylated to fluo-4, and fluo-4 can bind to intracellular Ca^{2+} to increase green fluorescence, so intracellular Ca^{2+} levels were detected by fluo-4/AM. After treatment with ADE, cells were incubated with 1 ml fluo-4/AM solution (10 mg/ml) for 30 min at 37°C. Microscopic images were obtained by a confocal microscopy (Nikon C2, Tokyo, Japan). ImageJ software was used to measure the fluorescence intensity.

2.9. Intracellular ROS Measurement. ROS level was assessed by DCFH-DA staining according to previous methods [30].

2.10. ER Structure Observation by Transmission Electron Microscopy (TEM). After being treated with ADE (1.25 mM) for 24 h, HT22 cells were digested with trypsin and fixed with 2.5% glutaraldehyde. Then, ultrathin sections (1 μm) were observed by TEM (Hitachi, Tokyo, Japan).

2.11. Quantitative Real-Time PCR (RT-PCR). RT-PCR was performed following a routine procedure. Total RNA was extracted from HT22 cells with TRIzol reagent according to the manufacturer’s instructions. The primer sequences were as follows.

(i) GRP78: 5′-GAAAGGAGGTGTGGGACG-3′ and 5′-CGCATCGCAATCAGACG-3′

(ii) PERK: 5′-CCGCAAGAAGGCCCTATCC-3′ and 5′-GAGTTTCAGACTCTTCCGCT-3′

(iii) ATF6: 5′-ACCTGTTCTCCTCTGAAATCCAA-3′ and 5′-AGGACAGAGAAAACAGCTGG-3′

(iv) Caspase-12: 5′-TGGAGGTTAGGCCAGACT-3′ and 5′-ATAGTGGGCATCTGGGTCTC-3′

(v) CHOP: 5′-AGTCCCTGCTCCTACCCT-3′ and 5′-GCTTTGGGATGTGCGTGTG-3′

(vi) GAPDH: 5′-GTATGACTCCACTCCAGGCAA-3′ and 5′-CACCAGTAGACTCCACGGACA-3′

2.12. Immunofluorescence Staining. The primary cultured neuron cells were seeded into six-well plates with cover glasses in each well and then treated with ADE and medium for 24 h. Immunofluorescence analysis was performed as previously described to determine βIII-tubulin and CHOP [30].

2.13. Western Blotting Analysis. After collecting ADE-treated HT22 cells, the proteins of the cells were extracted and the protein contents of the isolated supernatant were determined by a protein assay reagent. Equivalent amounts of protein (40 mg) were separated by gel electrophoresis. Then, the protein was transferred to a polyvinylidene difluoride (PVDF) membrane (Millipore Bedford, MA, USA). The membranes were immersed in 5% skimmed milk. Then, the primary antibodies were incubated with target antigens at 4°C overnight. The next day, horseradish peroxidase- (HRP-) conjugated secondary antibodies (1:3000) were incubated with the target antigens for 2 h. The enhanced chemiluminescence was used to visualize the proteins, and ImageJ software was used to do the densitometry analysis.

2.14. Statistical Analysis. Data were expressed as the mean ± SD of at least three representative experiments. Statistical significance was assessed using SPSS software 19.0 by one-way analysis of variance (ANOVA), followed by Dunnett’s multiple comparison tests. A p value of <0.05 was considered statistically significant.

3. Results

3.1. The Cytotoxicity of ADE. The MTT assay was used to evaluate the effect of ADE (0, 0.625, 1.25, 2.5, and 5 mM) on the cellular viabilities of HT22 cells. As shown in Figure 1(a), ADE exhibited cytotoxicity against HT22 cells in a dose-dependent manner. The cellular viabilities were decreased significantly after being exposed to ADE. As shown in Figure 1(b), the morphology of the HT22 cells has changed significantly characterized by cell shrinkage and irregular cell shapes. To further evaluate the effect of ADE-induced neurotoxicity, primary cultured cortical neuronal cells were treated with ADE (0, 1.25, 2.5, and 5 mM). As shown in Figure 1(c), ADE exhibited cytotoxicity against primary cultured cortical neuronal cells and the cellular viabilities were decreased significantly and dose-dependently after being exposed to ADE. As shown in Figure 1(d), primary cultured cortical neuronal cells exhibited obvious morphological changes characterized by shrinkage of neuronal cell bodies, reduction in the cell number and dendrites, and irregular cell shapes.

3.2. ADE Induces Apoptosis of HT22 Cells. HT22 cells were used to study whether ADE can induce apoptosis and the underlying mechanisms. As shown in Figure 2(a), after incubation with the ADE (0, 0.625, 1.25, and 2.5 mM) for 24 h, the morphological changes characterized by the nucleus pyknosis and fragmentation can be observed in
HT22 cells by Hoechst 33258 staining, indicating the appearance of apoptosis.

AO/EB staining was used to further distinguish the apoptosis of the cells. AO is a dye that penetrates normal cell membranes and causes cells to appear green in color. EB is a dye that passes through damaged cell membranes to cause cells to exhibit yellow fluorescence, demonstrating early apoptosis in cells. In our study, the nuclei of some cells appeared yellow or orange fluorescence after ADE treatment in Figure 2(b), which indicates that ADE disrupted the integrity of the cell membrane and might induce apoptosis in HT22 cells.

To clarify whether ADE induces apoptosis in HT22 cells, Annexin V-FITC/PI double-staining was used to measure the percentage of apoptotic cells in Figures 2(c) and 2(d). When compared with that in the control group, ADE treatment dose-dependently increased apoptotic ratio in HT22 cells. Therefore, these results suggest that ADE inhibited cell viabilities and induced apoptosis in HT22 cells.

Apoptosis-related proteins were further investigated to prove the ADE induced apoptosis. The data showed that the cleaved caspase-3, cleaved caspase-9, and cleaved caspase-12 expression levels were increased after ADE treatment in Figures 2(e) and 2(f). Treatment of HT22 cells with...
ADE also decreased intracellular Bcl-2 expression and increased Bax expression in Figures 2(e) and 2(f).

3.3. ADE Induces the ERS. In order to observe the changes of ER, TEM was used to directly observe the ultrastructural changes of ER after treatment with ADE (1.25 mM) for 24 h in HT22 cells. As shown in Figure 3(a), after treatment with ADE, the ER became swollen and vacuolated compared with the control group. In order to further prove the role of ERS in ADE-induced damage of HT22 cells, the expression levels of ERS-related mRNA were tested by RT-PCR. The data proved that the mRNA levels of GRP78, PERK, ATF6, caspase-12, and CHOP were dose-dependently upregulated in Figure 3(b).

Western blot was used to further demonstrate whether ERS is involved in cell apoptosis caused by ADE. As revealed in Figures 3(c) and 3(d), the protein expression levels of GRP78, ATF6, IRE1, p-IRE1, PERK, p-PERK, and CHOP were dose-dependently upregulated in the cells treatment with ADE. ERS-related CHOP protein was further detected

Figure 2: The exposure of ADE induces morphological changes of the HT22 cells. HT22 cells were incubated with ADE (0, 0.625, 1.25, and 2.5 mM) for 24 h, and then stained with Hoechst 33258 (a) and AO-EB (b), respectively. Scale bar = 50 μm. (c) Annexin V-FITC/PI double-staining of cells with ADE (0, 0.625, 1.25, and 2.5 mM) treatment for 24 h. (d) Quantified apoptotic rate (n = 3). (e) The expression levels of caspase family proteins, Bax and Bcl-2, by western blot. (f) Densitometric analysis of apoptosis-related proteins (n = 3, *p < 0.05, **p < 0.01 vs the control group). Statistical comparisons were made using one-way ANOVA with Dunnett’s multiple comparison tests.
by immuno-fluorescence in primary cultured cortical neuronal cells. Our data showed that ADE upregulated CHOP expression in a dose-dependent manner in Figure 3(e).

### 3.4. ERS Contributes to ADE-Induced Overproduction of Intracellular Ca²⁺ and ROS.

To further study the underlying mechanism of ADE-induced neurotoxicity, we tested the levels of ROS and Ca²⁺ in HT22 cells treated with ADE (0, 0.625, 1.25, and 2.5 mM) under a confocal microscope. The results showed that ADE treatment significantly increased intracellular ROS levels, indicating that ADE can induce oxidative stress in HT22 cells in Figures 4(a) and 4(b). Our data also demonstrated that ADE could significantly increase the intracellular Ca²⁺ levels Figures 4(c) and 4(d).

### 3.5. 4-PBA Attenuates ADE-Induced ERS and Apoptosis.

To verify the role of ERS in HT22 cell apoptosis, HT22 cells were treated with ERS inhibitor 4-PBA (1.25 mM) for 2 h before treatment with ADE (1.25 mM) for 24 h. The MTT assay proved that 4-PBA mitigated ADE-induced cell viability decrease in Figure 5(a). The swollen and vacuolated ER induced by ADE was improved after treatment with 4-PBA (Figure 5(b)). 4-PBA pretreatment also reduced ADE-induced increase of CHOP, GRP78, ATF6, cleaved caspase-9, cleaved caspase-12, and cleaved caspase-3 levels in Figures 5(c) and 5(d).

### 3.6. NAC Attenuates ADE-Induced ERS and Apoptosis.

Next, antioxidant NAC was used to further prove the role of oxidative stress in ERS-induced apoptosis. After pretreatment with NAC for 2 h, HT22 cells were treated with ADE (1.25 mM) for 24 h. As shown in Figure 6(a), the swollen and vacuolated ER induced by ADE was improved after pretreatment with NAC. We also proved that NAC reduced the accumulation
of ROS induced by ADE in HT22 cells in Figures 6(b) and 6(c). NAC pretreatment also reduced ADE-induced increase of CHOP, GRP78, ATF6, cleaved caspase-9, cleaved caspase-12, and cleaved caspase-3 levels in Figures 6(d) and 6(e).

4. Discussion

Excessive accumulation of ADE in the brain could lead to neurotoxicity and perhaps promote the development of neurodegenerative diseases [18]; however, the exact molecular mechanisms underlying ADE-induced neurotoxicity are not fully understood. Here, we studied the cytotoxic effects of ADE on HT22 cells and primary cultured cortical cells and found that ADE induced cytotoxicity by promoting apoptosis and inhibiting cell survival pathways.

Apoptosis is a programmed cell death that is usually caused by physiological or pathological factors. Evidence has shown that ADE is the most toxic metabolite of alcohol [4]. In various cell types, ADE is found to induce cytotoxicity and apoptosis [31, 32]. In this study, the destruction of the cell membrane integrity was demonstrated by staining the cells with AO/EB and Hoechst 33258. ADE was shown to induce apoptosis morphologically. Meanwhile, incubation with ADE caused obvious and dose-dependent increase of the percentage of dying cells by Annexin V-FITC/PI double-staining in HT22 cells, which was further proved in the primary cultured cortical cells. These results indicate that...
ADE induced cytotoxicity by inhibiting cell survival and promoting apoptosis.

Studies have shown that treatment with ADE leads to increase of intracellular ROS and oxidative stress in vivo [33]. ROS have close relationship with inflammation, oxidative stress, and cell damages [34, 35]. Previous investigations have shown that ADE can induce cytotoxicity by increasing the ROS accumulation and destroying redox homeostasis [3, 7]. In the current study, intracellular ROS levels were significantly increased by ADE treatment, suggesting that ADE can induce the oxidative stress by disrupting redox homeostasis in HT22 cells.

Ca²⁺ is an important second messenger that can participate in neurotransmitter release and signal transduction. Intracellular Ca²⁺ concentration changes can give rise to many cellular responses such as cell instability and cell damage [36]. At present, there is no literature on the relationship between ADE and Ca²⁺. To explore the possible mechanism of ADE-induced HT22 cell injury, intracellular Ca²⁺ levels after ADE treatment were examined. The results demonstrated that ADE can significantly increase intracellular Ca²⁺ concentration, which might be related to the disrupted redox homeostasis and ultimately contribute to ADE-induced apoptosis in HT22 cells.

ER is an important organ regulating the intracellular environmental stability and is susceptible to oxidative stress [37]. In addition, ERS is closely related to the destruction of intracellular Ca²⁺ homeostasis. On the other hand, Ca²⁺ can...
regulate the expression of ER molecular chaperones [38]. Therefore, we believed that the increased levels of intracellular ROS and Ca^{2+} might influence the function of ER (Figure 7).

Recent evidence indicates that ERS may play a role in the pathogenesis of neurological diseases [39] and has been implicated in various neurodegenerative processes, such as brain ischemia [40], Alzheimer’s disease [41], Parkinson’s disease [42], and amyotrophic lateral sclerosis [43]. The role of ER in the apoptotic process has been well established. ER apoptotic pathway is mainly regulated by activating GRP78, upregulating the expression of CHOP and cleave caspase-12, and inducing apoptosis [44]. The activation of ERS response or UPR is mediated mainly by 3 transmembrane ER signaling proteins: PERK, IRE1, and ATF6. In the current study, the mRNA levels of GRP78, PERK, ATF6,
CHOP, and caspase-12 were dose-dependently upregulated after ADE treatment. Furthermore, ERS-related proteins including IRE1, p-IRE1, ATF6, PERK, p-PERK, GRP78, and CHOP tested by western blot were also significantly upregulated after ADE treatment. The increase of PERK mRNA level is much more robust than the change of PERK protein, because PERK protein has been significantly phosphorylated to p-PERK after ADE treatment. Both PERK and IRE1 are type I transmembrane proteins of ER. PERK and IRE1 dissociate from the chaperone protein GRP78 when occurring ERS, forming phosphorylated forms of p-PERK and p-IRE1. In the present study, the protein levels of IRE1, p-IRE1, PERK, and p-PERK were all increased after ADE treatment, which strongly indicates the appearance of ERS.

To further examine ERS-mediated apoptosis, we tested the expression levels of CHOP and caspase-12. CHOP is a member of the bZIP transcription factor C/EBP family [39]. Under normal conditions, the expression level of CHOP is very low. When ERS occurs, the three main ERS pathways including IRE1, ATF6, and PERK can simultaneously stimulate CHOP [40, 41]. Caspase-12 is a unique endoplasmic reticulum apoptosis protease that can participate in ERS-induced apoptosis [42]. Our results suggest that following ADE treatment of HT22 cells, CHOP and cleaved caspase-12 levels were significantly increased, which were consistent with the changes at mRNA levels. We further measured CHOP protein in primary cultured cortical cells by immunofluorescence and showed that ADE obviously increased the expression of CHOP. These data indicate that ADE-induced apoptosis correlates with activated ERS sensors and upregulates ERS-related protein expression (Figure 7).

Basic and clinical studies over the past decade have shown that ROS have a significant impact on the pathogenesis of many human diseases [43]. In ERS, disorganized disulfide bond formation and disruption may lead to ROS accumulation and oxidative stress [44]. At the same time, ERS can cause mitochondrial damage and increase mitochondrial ROS production [45]. In addition, some UPR components such as CHOP may cause oxidative stress. In many in vitro and in vivo models related to ERS, ERS and oxidative stress emphasize each other in a positive feed-forward loop, which interferes with cell function and activates proapoptotic signaling [46]. In this study, the antioxidant NAC effectively inhibited the production of ROS and also alleviated the ERS (Figure 7). The results indicate that oxidative stress may promote ERS and induce cytotoxicity.

Here, for the first time, we proved that ADE induced neurotoxicity of HT22 cells via oxidative stress- and Ca\(^{2+}\) imbalance-mediated ERS. We explored the consequences of ADE on ER homeostasis and ROS production and found that ADE induced excessive generation of intracellular ROS and upregulation of ERS-related proteins GRP78, p-PERK, and p-IRE1. ER transmembrane proteins sense the folding state of proteins inside the ER lumen and convey the information to the mitochondria and the cytosol. 4-PBA, an ERS inhibitor, acts as a chemical chaperone assisting protein folding...
in the ER as well as in the cytosol and the mitochondria [47]. In our results, inhibition of ROS with antioxidant NAC and mitigation of ERS with ERS inhibitor 4-PBA obviously suppressed ADE-induced apoptosis and the expression of ERS-related proteins. Meantime, we proved that antioxidant NAC not only mitigated ADE-induced ROS increase but also suppressed the swollen and vacuolated ER caused by ADE. Notably, the abnormal levels of ERS caused by ADE were significantly alleviated when ROS was inhibited by NAC. Therefore, on the basis of these findings, we believe that the activation of excessive ROS is a pathway contributing to the generation of ERS caused by ADE in the HT22 cells.

5. Conclusion
We demonstrated for the first time that ADE can cause neurotoxicity in HT22 cells and primary cultured cortical neuronal cells. ADE induced apoptosis of HT22 cells via the promotion of ERS, and excessive ROS and Ca\(^{2+}\) were involved in ADE-induced ERS. Our study also provides evidence that ERS inhibitor 4-PBA and antioxidant NAC may be beneficial for preventing neuronal damage associated with ADE-induced cytotoxicity which could be resulting from excessive alcohol consumption (Figure 7). Our data provides new insights into the underlying mechanisms of ADE-induced neurotoxicity. The ERS induced by ADE may have a profound effect on the CNS, which requires further study.

Abbreviations
ADE: Acetaldehyde
ADH: Alcohol dehydrogenase
ALDH: Aldehyde dehydrogenase
ERS: Endoplasmic reticulum stress
NAC: N-acetyl-L-cysteine
ROS: Reactive oxygen species
4-PBA: Sodium 4-phenylbutyrate.

Data Availability
The data used to support the findings of this study are available from the corresponding authors upon request.

Conflicts of Interest
The authors declare that there is no conflict of interests regarding the publication of this paper.

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