Chronic ethanol exposure induces SK-N-SH cell apoptosis by increasing N-methyl-D-aspartic acid receptor expression and intracellular calcium

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Abstract. It has been identified that chronic ethanol exposure damages the nervous system, particularly neurons. There is scientific evidence suggesting that neuronal loss caused by chronic ethanol exposure has an association with neuron apoptosis and intracellular calcium oscillation is one of the primary inducers of apoptosis. Therefore, the present study aimed to investigate the inductive effects of intracellular calcium oscillation on apoptosis in SK-N-SH human neuroblastoma cells and the protective effects of the N-methyl-D-aspartic acid receptor (NMDAR) antagonist, memantine, on SK-N-SH cell apoptosis caused by chronic ethanol exposure. SK-N-SH cells were treated with 100 mM ethanol and memantine (4 µM) for 2 days. Protein expression of NR1 was downregulated by RNA interference (RNAi). Apoptosis was detected by Annexin V/propidium iodide (PI) double-staining and flow cytometry and cell viability was detected using an MTS kit. Fluorescence dual wavelength spectrophotometry was used to determine the intracellular calcium concentration and the levels of NR1 and caspase-3 were detected using western blotting. NR1 mRNA levels were also detected using qPCR. It was found that chronic ethanol exposure reduced neuronal cell viability and caused apoptosis of SK-N-SH cells, and the extent of damage in SK-N-SH cells was associated with ethanol exposure concentration and time. In addition, chronic ethanol exposure increased the concentration of intracellular calcium in SK-N-SH cells by inducing the expression of NMDAR, resulting in apoptosis, and memantine treatment reduced ethanol-induced cell apoptosis. The results of the present study indicate that the application of memantine may provide a novel strategy for the treatment of alcoholic dementia.

Introduction

Ethanol is eventually metabolized to carbon dioxide and water via a dehydrogenase and oxidase. Ethanol and a series of intermediate products in its metabolism have certain toxic effects on tissues and cells, which may result in a wide range of mental and physical abnormalities (1). Chronic alcoholism may damage multiple systems and organs including the nervous, cardiovascular, digestive and immune system, as well as muscle. In a previous study, it was demonstrated that ethanol may lead to the loss of neurons in specific brain regions (2). It has been suggested that cell necrosis or apoptosis may be activated by ethanol via a variety of mechanisms, such as enhancing expression of tumor necrosis factor-α (3,4) and Fas/Fas ligand (5,6), inducing transcriptional activity of nuclear factor-κB (7,8), activating caspases in mitochondria (9) and causing transport disorders of intracellular Ca²⁺ (10). Intracellular Ca²⁺ serves an important role in cell biology, which influences or determines necrosis or apoptosis independently. Cellular Ca²⁺ overload is the hub of necrosis or apoptosis (11,12). Intracellular Ca²⁺ originates from L-type voltage-operated calcium channels, N-methyl-D-aspartate receptor (NMDAR)-mediated calcium influx into cells (13), inositol 1,4,5-trisphosphate (IP3)-induced calcium release (14) and ryanodine receptor-mediated calcium release (15). A previous report has identified a rise of excitatory amino acids in patients with chronic alcohol poisoning (16), such as glutamate, aspartate and homocysteine, which significantly increased the vulnerability of neurons to excitotoxicity and oxidative damage. Excitatory amino acids serve a role in...
NMDAR overstimulation, oxidative stress, caspase activation, DNA damage and mitochondrial dysfunction (17). NMDAR is a ligand-gated Ca\(^{2+}\) channel that is closely associated with central nervous system development, learning and memory (18,19). Following chronic ethanol exposure, NMDAR in the central nervous system is more sensitive to NMDA, which is known as NMDAR hyper-excitement (20), a primary cause of ethanol withdrawal symptoms and neuronal over-excitability (21). NMDAR is heteromeric and consists of three subunits: NR1, NR2 and NR3 (22). The NR1 subunit is the functional subunit of NMDAR, which is widespread in the central nervous system of animals (23). It has been demonstrated previously that, following long-term ethanol feeding, cerebral cortical MK-801 binding in guinea pigs was significantly higher than that in a control group (24) and there appeared to be greater expression of NR1 in the cerebral cortex, hypothalamus and hippocampus of rats (25). These findings suggested that long-term ethanol exposure results in an adaptive increase in NMDAR, causing hyper-excitation of NMDAR. Preconditioning with non-competitive NMDAR antagonist, memantine and downregulation of NMDAR expression by small interfering RNA (siRNA) control intracellular Ca\(^{2+}\) release (26), thereby inhibiting caspase-3 activation and controlling or reducing the occurrence of neuronal apoptosis induced by isoflurane. The present study aimed to determine the role of NMDAR and intracellular calcium in ethanol-induced SK-N-SH cell apoptosis and assess the neuroprotective and therapeutic effect of memantine.

Materials and methods

Cell line. In the present study, SK-N-SH human neuroblastoma cells were purchased from Nanjing KeyGen Biotech Co., Ltd. (Nanjing, China). The cells were cultured in high glucose Dulbecco's modified Eagle's medium (DMEM; Biological Industries, Kibbutz Beit-Haemek, Israel) containing 10% fetal bovine serum, 100 U/ml penicillin and 100 µg/ml streptomycin (all purchased from Biological Industries). Cells were maintained at 37°C in a humidified atmosphere containing 5% CO\(_2\).

Ethanol volatilization detection. The present study determined that ethanol volatilized over time under normal culture conditions, thereby reducing the ethanol concentration in the culture medium. Therefore, ethanol volatilization was determined per 24 h and appropriate quantities of ethanol were added to maintain relatively stable concentrations of ethanol in the medium. 5 ml culture medium containing ethanol (50, 100, 200 and 400 mM) was added into a 25-cm\(^2\) cell culture flask. After 24 h at 37°C, the remaining ethanol concentration was detected using headspace gas chromatography (cat. no. GC-14A; Shimadzu Corporation, Kyoto, Japan), as described in a previous study (27). The quantity of daily ethanol volatilization was calculated using the following equation: Quantity of daily ethanol volatilization = initial quality - remaining quality.

Grouping. SK-N-SH cells were cultured in a 25-cm\(^2\) cell culture flask at a density of 1x10\(^5\) cells/ml. Ethanol was added into DMEM culture medium at 37°C. When cells reached an 80-90% confluence, according to the duration of ethanol treatment, SK-N-SH cells were divided into 24, 48 and 72 h groups, and according to the ethanol concentration, cells were divided into 0 (control group), 50, 100, 200 and 400 mM groups. Ethanol at 0 and 100 mM was also used to treat SK-N-SH cells for 2 days at 37°C and cells were categorized into memantine (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) and non-memantine groups. The concentration of memantine used was 4 µM. According to whether protein expression of NR1 was downregulated by RNA interference (RNAi), cells were divided into NR1 short hairpin RNA (shRNA) and control shRNA groups.

RNAi. SK-N-SH cells were seeded in 6-well plates at a low density (<10% confluence) in normal growth medium containing various concentrations of puromycin (cat. no. sc108071; Santa Cruz Biotechnology, Inc., Dallas, TX, USA) (0, 1, 2, 3, 4, 6, 8 and 10 µg/ml) to determine the minimum concentration necessary to kill all untransfected cells. An NR1 shRNA plasmid (cat. no. sc-91941-SH; Santa Cruz Biotechnology, Inc.) was used in the RNAi. It is a pool of 3 target-specific lentiviral vector plasmids, each encoding 19-25 nucleotide (plus hairpin) shRNAs designed to knock down gene expression. The control shRNA plasmid (cat. no. sc-108060; Santa Cruz Biotechnology, Inc.) encodes a scrambled shRNA sequence that does not degrade any known cellular mRNA. The NR1 shRNA plasmid and the control shRNA plasmid were of the same vector type. Each plasmid contained a puromycin resistance gene for the selection of stable cells that express the desired shRNA. At 60-70% confluence, shRNA Plasmid Transfection Reagent (10/200 µl, cat. no. sc-108061; Santa Cruz Biotechnology, Inc.) was used to transfect the NR1 shRNA plasmid (5/200 µl) and control shRNA plasmid (5/200 µl) into cells, according to the manufacturers’ protocol. Following 48 h transfection at 37°C, the medium was replaced with fresh DMEM medium containing puromycin (1 µg/ml) to select the transfected cells over 5 days at 37°C. At 90% confluence, total protein was extracted and the transfection efficiency was determined by western blotting. The sequences of NR1 shRNA and control shRNA (28) are presented in Table I.

MTS assay. Cell viability was determined using an MTS kit (Promega Corporation, Madison, WI, USA). SK-N-SH cells were seeded at a density of 4,000 cells/well and treated in 96-well plates for various durations. Following 24 h culture, the cells were washed with PBS three times and fresh DMEM medium (100 µl) and MTS (20 µl) reagent were added to the wells for 1 h at 37°C in the dark (25,26). The absorbance was measured at a wavelength of 490 nm on an ELx808 absorbance reader (BioTek Instruments, Inc., Winooski, VT, USA). To eliminate possible interference by alcohol, cells treated with the same concentrations of alcohol (0, 50, 100, 200 and 400 mM) and memantine (4 µM) but without addition of assay reagents were used as blank controls. All experiments were repeated at least five times.

Annexin V/propidium iodide (PI) double-staining. Cell apoptosis was measured using an Annexin V-fluorescein isothiocyanate (FITC)/PI apoptosis detection kit (BD Biosciences, San Jose, CA, USA). SK-N-SH cells were seeded at a density of 1x10\(^5\) cells/ml and treated in 25-cm\(^2\) tissue culture flasks until an 80-90% confluence was observed. Following washing with PBS twice, cells were double-stained with FITC-conjugated Annexin V and PI.
for 15 min at 20˚C in a Ca\(^{2+}\)-enriched binding buffer in the kit. Cells were immediately analyzed on a flow cytometer in their staining solution. Annexin V and PI emissions were detected in the FL‑1 (band pass, 530 nm; band width, 30 nm) and FL‑2 (band pass, 585 nm; band width, 42 nm) channels. A total of 10,000‑20,000 events were recorded per sample. BD FACSDiva V8.0.1 software (BD Biosciences) was used to analyze this data.

**Intracellular calcium measurement.** Intracellular Ca\(^{2+}\) was measured with the Ca\(^{2+}\)-sensitive dye fura‑2‑acetoxymethyl ester (fura‑2‑AM; Dojindo Molecular Technologies, Inc., Kumamoto, Japan) as previously described (29). SK‑N‑SH cells were seeded to 80‑90% confluence and treated in 25‑cm\(^2\) tissue culture flasks. To prepare cell suspensions, the cells were washed twice with Hank's balanced salt solution (HBSS; Biological Industries), trypsinized (Biological Industries), centrifuged at a speed of 2,000 x g, for 5 min at room temperature) and resuspended in HBSS containing 20 g/l bovine serum albumin (Biological Industries), and the cell concentration was adjusted to 1x10^6‑1x10^7 cells/ml. The survival rate of the cells was determined using a trypan blue staining cell viability assay kit (Beyotime Institute of Biotechnology, Shanghai, China) according to the manufactures' protocol. A total of 0.1 ml cell suspension, containing 10^5 cells was mixed with 0.4% trypan blue solution (0.1 ml). The mixture was then incubated for 3 min at room temperature. The quantity of living cells (which were not stained by trypan blue) were counted and determined to be >95%. SK‑N‑SH cells (1x10^6‑3x10^6 cells/ml) were loaded with 3 mM fura‑2‑AM in HBSS at 37˚C for 20 min. The cells were then washed once with HBSS and incubated for 1 h at 37˚C in HBSS. The fluorescence of the cell suspension was monitored continuously using a F‑4500 fluorescence spectrometer (Hitachi, Ltd., Tokyo, Japan) with excitation at 340 and 380 nm and emission at 500 nm. Triton X‑100 (0.1%) and 10 mM EDTA were added to obtain the maximum and minimum concentrations of calcium, respectively.

**Cell lysis and protein quantification.** SK‑N‑SH cells were seeded until an 80‑90% confluence was reached and treated with ethanol (0, 50, 100, 200 and 400 mM) and memantine (4 µM) in 25‑cm\(^2\) tissue culture flasks. Cells were subsequently washed with Dulbecco's PBS (Biological Industries) and lysed on ice using radioimmunoprecipitation assay lysis buffer (Beyotime Institute of Biotechnology) containing 1 mM phenylmethanesulfonyl fluoride (Beyotime Institute of Biotechnology). Lysates were harvested using cell scrapers, placed on ice for 30 min and then fragmented using an ultrasonicator. The lysates were centrifuged at 21,000 x g for 15 min at 4˚C and quantified for total proteins using a bicinchoninic acid protein assay kit (Beyotime Institute of Biotechnology).

**Western blotting.** Equal quantities of protein (~30 µg) were separated using 10% SDS‑PAGE and the separated protein was transferred onto polyvinylidene fluoride membranes. The membranes were blocked for non‑specific binding with 5% non‑fat dry milk in Tris-buffered saline containing 0.05% Tween‑20 (TBS‑T) for 2 h at room temperature and then probed with primary antibodies overnight at 4˚C. The primary antibodies used were as follows: Rabbit anti‑NR1 (1:500; cat. no. 13771‑1‑AP; ProteinTech Group, Inc., Chicago, IL, USA), rabbit anti‑caspase‑3 (1:200; cat. no. sc‑98785; Santa Cruz Biotechnology, Inc.) and mouse anti‑β‑actin (1:1,000; cat. no. TA‑09; OriGene

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**Table I. Sequences of shRNA.**

| shRNA                | Sequence                                                                 |
|----------------------|--------------------------------------------------------------------------|
| NR1 shRNA‑A          | Hairpin sequence 5’GATCCCCATGTTCTTAGAGAAGATTTCAAGAGAATCTTTCTTAAGAAACATGTTTTT’3 |
|                      | Corresponding siRNA sense sequence 5’CCAUGUUUCUUAGAGAAGAUtr3               |
|                      | Corresponding siRNA antisense sequence 5’AUCUUUCUAAAGAAACUGGtr3            |
| NR1 shRNA‑B          | Hairpin sequence 5’GATCCCTTGTATTTGTCGGGAAGATTTCAAGAGATCTTTCTCCGACAATACAAGTTTTT’3 |
|                      | Corresponding siRNA sense sequence 5’CUUGUAUUGUGCGGAAAGtr3                |
|                      | Corresponding siRNA antisense sequence 5’UCUUUCCCGACAAUACAAAGtr3           |
| NR1 shRNA‑C          | Hairpin sequence 5’GATCCCAAGGTGGATCCAGTCTTTCTTAAGAGAAGAAACTGGATCCACCTTGTTTTT’3 |
|                      | Corresponding siRNA sense sequence 5’CAAGGGUGGAUCCAGUUCUtr3                |
|                      | Corresponding siRNA antisense sequence 5’AGAAACUGGAUCCACCUUGtr3            |
| Control shRNA        | 5’TTCTCCGAACGTCACGTTCACAGTTCTCAAGAGAACGTGACACGTTCGGAGAATTTTT’3             |

shRNA, short hairpin RNA.
Technologies, Inc., Rockville, MD, USA). Subsequently, the blots were washed with TBS-T three times (5 mins/wash) and incubated with the corresponding peroxidase-conjugated goat anti-mouse or anti-rabbit secondary antibodies (1:10,000; cat. nos. E00310-02 and E00312-02; EarthOx Life Sciences, Millbrae, CA, USA) for 2 h at room temperature. Protein bands were detected with enhanced chemiluminescence reagent (EMD Millipore, Billerica, MA, Germany). Chemiluminescent signals were detected and analyzed by a Tanon 5500 Chemiluminescent Imaging system (Tanon Science and Technology Co., Ltd., Shanghai, China). Band densities were analyzed semi-quantitatively using Image J 1.6.0 software (National Institute of Health, Bethesda, MD, USA).

**RNA extraction and reverse transcription-quantitative polymerase chain reaction (RT-qPCR).** Total RNA was extracted from cells using TRIzol® (Thermo Fisher Scientific, Inc., Waltham, MA, USA) and reverse transcribed into cDNA using a PrimeScript™ RT Reagent kit (Perfect Real Time; Takara Biotechnology Co., Ltd., Dalian, China). qPCR was performed using a SYBR® Premix Ex Taq™ II (TliRnaseH Plus; Takara Biotechnology Co., Ltd.) in a reaction volume of 20 µl on an ABI 7500 Real-Time PCR system (Thermo Fisher Scientific, Inc.) using the following thermocycling conditions: 95˚C for 30 sec, followed by 40 cycles of 95˚C for 5 sec and 60˚C for 34 sec, 95˚C for 15 sec, 60˚C for 60 sec and 95˚C for 15 sec. The primer sequences used were as follows: β-actin forward, 5'-CTAACTTGGCAGAAAAACAAGAT-3' and reverse, 5'-TTCTGTGAAACGCATCTCTCA-3'; and NMDAR1 forward, 5'-CGCCAACACAGCATCATT-3' and reverse, 5'-ATCTGTCACAATCTTCTAGTCT-3'. β-actin was used as the reference gene. The relative gene expression levels were represented as ΔΔ quantification cycle (ΔΔCq) and the fold change of gene expression was calculated via the 2^ΔΔCq method (30). Experiments were repeated in triplicate.

**Statistical analysis.** GraphPad Prism version 6 (GraphPad Software, Inc., La Jolla, CA, USA) was used for statistical analysis. Measurement data are expressed as the mean ± standard error. One-way analysis of variance and Turkey's multiple comparisons test was used to compare differences between groups. The Wilcoxon rank sum test was used to compare differences among other types of data, including the percentage of apoptotic cells. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**Daily ethanol volatilization.** The ethanol volatilization per 24 h from 25-cm² flasks containing 5 ml culture medium with ethanol are presented in the Table II.

| Groups          | % daily ethanol volatilization (mean ± standard error) |
|-----------------|--------------------------------------------------------|
| 50 mM ethanol   | 19.57±1.37                                              |
| 100 mM ethanol  | 22.12±1.07                                              |
| 200 mM ethanol  | 24.57±0.60                                              |
| 400 mM ethanol  | 27.55±1.01                                              |

Figure 1. NR1 expression in SK-N-SH cells following chronic ethanol exposure. (A and B) Relative expression of NR1 in SK-N-SH cells is presented. All data were normalized to expression levels of β-actin. The relative expression levels of the NR1 protein in the control shRNA group was similar to the control group and the difference was not statistically significant. Conversely, the relative expression levels of the NR1 protein in the NR1 shRNA group was significantly lower than that in the control shRNA group. *P<0.01 vs. control shRNA group. Data are presented as the mean ± standard error of the mean (n=6). NR1, N-methyl-D-aspartate receptor subunit 1; shRNA, short hairpin RNA.

**SK-N-SH cell activity decreases following chronic ethanol exposure and the effect is attenuated by memantine and downregulation of NR1 protein.** SK-N-SH cells were treated with increasing concentrations of ethanol for 24-72 h and then cell viability was measured. Compared with the 24 h/0 mM ethanol group, cell viability decreased with increasing ethanol concentration and time (Fig. 2A). Compared with the control group, the cell viability of the ethanol groups was significantly lower. Compared with the ethanol group, the cell viability of the ethanol + memantine and ethanol + NR1 shRNA groups was significantly higher (Fig. 2B).

**Apoptosis of SK-N-SH cells increases following chronic ethanol exposure and the effect is attenuated by memantine and downregulation of the NR1 protein.** SK-N-SH cells were treated with increasing concentrations of ethanol for 24-72 h.
Annexin V-FITC/PI double staining was used to detect the apoptotic rate of cells. Apoptotic cells included early apoptotic and late apoptotic cells (Fig. 3A and B; Table III). Compared with the 0 mM ethanol groups at 24, 48 and 72 h, apoptosis was increased gradually with increasing ethanol concentration and time (Fig. 3A and B). Compared with the control group, apoptosis of the ethanol group was significantly higher. Compared with the ethanol group, apoptosis of the ethanol + memantine and ethanol + NR1 shRNA groups were significantly lower (Fig. 3C and D).

Memantine and downregulation of NR1 protein attenuates activation of the caspase-3 protein induced by ethanol. Western blotting revealed that the expression of cleaved caspase-3 was higher in the ethanol group compared with the control group, whereas expression of cleaved caspase-3 was significantly lower in the ethanol + memantine and ethanol + NR1 shRNA groups compared with the ethanol group (Fig. 4A and B).

Ca^{2+} concentrations in SK-N-SH cells increase following chronic ethanol exposure and the effect is attenuated by memantine and downregulation of the NR1 protein. Compared with the 0 mM ethanol groups at 24-72 h, the mean intracellular Ca^{2+} concentration increased with increasing ethanol exposure concentration and time (Fig. 5A). Compared with the control group, the mean intracellular calcium concentration in the ethanol group was significantly higher. Compared with the ethanol group, the mean intracellular calcium concentration in the ethanol + memantine and ethanol + NR1 shRNA groups was significantly lower (Fig. 5B).

NR1 protein and mRNA expression levels in SK-N-SH cells increase following chronic ethanol exposure. SK-N-SH cells were treated with increasing concentrations of ethanol for 24-72 h. Whole proteins were extracted for western blotting and the relative expression levels of the NR1 protein was detected. β-actin was used as the internal reference. Compared with the 0 mM ethanol groups at 24-72 h, relative expression levels of the NR1 protein increased with increasing ethanol concentration and time (Fig. 6A-F). SK-N-SH cells were treated with increasing concentrations of ethanol for 24-72 h, and then the relative expression levels of NR1 mRNA were measured via RT-qPCR. Fig. 6G-I demonstrate that, compared with the 0 mM ethanol groups at 24, 48 and 72 h, the relative mRNA expression of NR1 increased gradually (excluding the 50 mM group at 24 h) with increasing ethanol concentration and time (P<0.05; Fig. 6G-I).

### Table III. Analysis of apoptosis in each group.

| Groups                  | % cell apoptosis (mean ± standard error) |
|-------------------------|----------------------------------------|
| 24 h                    |                                        |
| Control                 | 2.66±0.19                              |
| 50 mM ethanol           | 3.72±0.13                              |
| 100 mM ethanol          | 4.60±0.25                              |
| 200 mM ethanol          | 12.54±0.58*                            |
| 400 mM ethanol          | 13.96±0.70*                            |
| 48 h                    |                                        |
| Control                 | 3.72±0.30                              |
| Memantine               | 3.22±0.27                              |
| NR1 shRNA               | 4.50±0.25                              |
| 50 mM ethanol           | 4.36±0.42                              |
| 100 mM ethanol          | 15.06±1.08*                            |
| 100 mM ethanol + memantine | 7.02±0.35*                            |
| 100 mM ethanol + NR1 shRNA | 9.42±0.48*                            |
| 200 mM ethanol          | 25.04±1.18*                            |
| 400 mM ethanol          | 30.94±1.28*                            |
| 72 h                    |                                        |
| Control                 | 3.92±0.55                              |
| 50 mM ethanol           | 7.04±0.51                              |
| 100 mM ethanol          | 22.00±1.59*                            |
| 200 mM ethanol          | 31.26±1.26*                            |
| 400 mM ethanol          | 38.32±1.62*                            |

*P<0.01 vs. control group; ^P<0.01 vs. 100 mM ethanol group.
Discussion

Previous studies have indicated that light, moderate to chronic or acute ethanol consumption may reduce neuron death and exhibit potentially neuroprotective effects (31,32). But most studies have reported nerve cell degeneration, apoptosis and reduced densities in deceased individuals who succumb to chronic ethanol poisoning, as well as in experimental animals and cultured cells that undergo chronic ethanol exposure (33,34). These findings indicated that ethanol induces death.
nerve cell apoptosis via a number of mechanisms (35). It has previously been speculated that ethanol causes neuronal ATP metabolic disorders and calcium overload by decreasing cytochrome oxidase activity (36). Ethanol under the action of dehydrogenase and aldehyde dehydrogenase produce oxygen-free radicals that damage nerve cell DNA (37). As such, there is a strong interest in studies assessing ethanol neurotoxicity. However, due to the self-repair of cells and volatility of ethanol, the underlying mechanism of chronic ethanol exposure during apoptosis of cultured neurons in vitro remains unclear (38,39). A previous study has suggested that the use of siRNAs to downregulate gene expression of NMDAR, IP3 receptor or sarco/endoplasmic reticulum calcium adenosine triphosphatase ATP-1 (SERCA1), or pre-administration of non-competitive NMDAR antagonist, memantine, inhibit intracellular Ca\(^{2+}\) release, thereby inhibiting the activation of caspase-3 induced by isoflurane to control and reduce the occurrence of neuronal apoptosis (26). To the best of our knowledge, no previous studies have assessed the relationship between ethanol, NMDAR, intracellular Ca\(^{2+}\) and apoptosis.

In the present study, SK-N-SH human neuroblastoma cells were used to examine whether ethanol-induced apoptosis was associated with NMDAR and intracellular calcium. SK-N-SH cells have been used in a previous study of neuronal cell apoptosis (40). Ethanol has strong volatility and many in vitro studies of ethanol exposure have investigated the effect of ethanol on cells cultured for a short period of time (41,42). In a preliminary experiment, it was observed that due to its strong volatility, ethanol is unable to maintain relatively stable concentrations, which is accompanied by a compensatory response of self-protection by SK-N-SH cells. Therefore, the present study performed an ethanol volatilization experiment to ensure maintenance of chronic ethanol exposure.

Compared with the control group, an increase was observed in NR1 protein expression, mean intracellular Ca\(^{2+}\) concentration and apoptotic rate of SK-N-SH cells, and a decrease was observed in cell viability. It was also demonstrated that with greater exposure concentration and duration of ethanol to SK-N-SH cells, the degree of cell damage was increased. These results indicated successful establishment of the chronic ethanol exposure model in SK-N-SH cells and confirmed that expression of the NR1 protein in SK-N-SH cells was increased by chronic ethanol exposure. As the expression of NMDAR increases, the resting intracellular Ca\(^{2+}\) concentration in the cell membrane increases, thereby increasing the ability of the cell to resist apoptosis.
NR1 protein increased, the intracellular calcium concentration also increased. This suggested that the effects of chronic ethanol exposure may be mediated via the NMDAR-mediated calcium transport pathway to increase the intracellular calcium concentration in SK-N-SH cells. High intracellular calcium may activate apoptosis and reduce cell viability and proliferation.

To support the above speculation, SK-N-SH cells were treated with 100 mM ethanol for 48 h and then with the noncompetitive NMDAR antagonist, memantine, at 4 µM. In addition, the expression levels of the NR1 gene in SK-N-SH cells was downregulated by NR1 shRNA. The results revealed an increase in the mean Ca^{2+} concentration, of cleaved caspase-3 and apoptosis, and a decrease in cell viability of the ethanol group compared with the control group. Compared with the ethanol group, there were decreases in the mean intracellular Ca^{2+} concentration, expression of cleaved caspase-3 and apoptotic rate, and an increase in the cell viability of the ethanol + memantine and ethanol + NR1 shRNA groups.

However, the present study also observed that shRNA-mediated downregulation of NR1 protein expression and non-competitive antagonism by memantine did not completely reverse the increase in the intracellular Ca^{2+} concentration, increase in apoptosis, or the decrease of cell viability and other neurotoxic effects caused by chronic ethanol exposure in neuronal cells. These results suggested that the neurotoxicity caused by chronic ethanol exposure is not limited to its effect on NMDAR, but also involves a variety of mechanisms working together. The mechanism of cellular damage caused by chronic ethanol exposure to neuronal cells is complex. Therefore, a follow-up study of IP3R (43), SERCA1 (44) and other proteins associated with Ca^{2+} transport pathway and other signaling pathways associated with neuronal cell damage, will be investigated in future studies.

In conclusion, chronic ethanol exposure inhibited neuronal cell viability and caused apoptosis of neuronal SK-N-SH cells, and the extent of damage in SK-N-SH cells was associated with ethanol exposure concentration and duration. In addition, chronic ethanol exposure induced expression of NMDAR and increased the concentration of intracellular Ca^{2+} in SK-N-SH cells, resulting in apoptosis. Memantine had a protective effect against damage in SK-N-SH cells. The results of the present study indicate that the application of memantine may provide a novel strategy in the treatment of alcoholic dementia. However,
future studies should be conducted in vivo using animals to assess the effects of ethanol on the brain, including in learning and memory.

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Availability of data and materials

The analyzed data sets generated during the present study are available from the corresponding author on reasonable request.

Authors’ contributions

The work presented here was carried out in collaboration between all authors. GZ and RZ collaborated to design the study. HW, XW, YL, HY and JC the designed methods and experiments. HW, CW, GX and JY carried out the laboratory experiments. CF and PW analyzed the data. HW, XW and YL drafted the manuscript. XW, RZ and GZ provided critical revision and contributed to the interpretation of findings. All authors have contributed to, read and approved the manuscript.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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