Effects of Snake Venom Pharmacopuncture on a Mouse model of Cerebral Infarction

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ABSTRACT

Background: This study investigated the effects of Vipera lebetina turanica snake venom (SV) on cerebral infarction induced by middle cerebral artery occlusion in mice.

Methods: Following cerebral infarction, SV was injected intravenously or added to BV2 cell culture. Tissue injury was detected using triphenyltetrazolium chloride (TTC) staining, neurological deficit score, NO, ROS, and GSH/GSSG assays, qPCR, Western blot, and cell viability.

Results: Cerebral infarction caused by middle cerebral artery occlusion as observed by TTC staining, showed SV inhibited cell death, reducing the number of brain cells injured due to infarction. SV treatment for cerebral infarction showed a significant decrease in abnormal behavior, as determined by the neurological deficit score. The oxidation and inflammation of the cells that had cerebral infarction caused by middle cerebral artery occlusion (NO assay, ROS, GSH/GSSG assay, and qPCR), showed significant protection by SV. Western blot of brain infarction cells showed the expression of iNOS, COX-2, p-IkB-α, P38, p-JNK, p-ERK to be lower in the SV group. In addition, the expression of IkB increased. BV2 cells were viable when treated with SV at 20 μg/mL or less. Western blot of BV2 cells, treated with 0.625, 1.5, 2.5 μg/mL of SV, showed a significant decrease in the expression of p-IkB-α, p-JNK, iNOS, and COX-2 on BV2 cells induced by LPS.

Conclusion: SV showed anti-inflammatory and anti-oxidant effects against cerebral infarction and inflammation.

Introduction

Ischemic brain injury is a worldwide medical problem that often causes irreversible brain damage. Moreover, the prevention of ischemic brain injury is key since the after-effects of a stroke significantly jeopardizes quality of life. Several risk factors predispose cerebral infarction, e.g., chronic hypertension, diabetes, smoking, and hypercholesteremia, and prognoses are unfavorable in the presence of these risk factors [1].

Several studies have reported that inhibiting inflammatory processes that produce cyclooxygenase (COX) -2 (COX-2) and inducible nitric oxide synthase (iNOS) reduce the number of strokes in animal models [2]. Many pro-inflammation enzymes such as iNOS, COXs and xanthine oxidase, participate in oxidative injury during cerebral ischemia [3]. iNOS is found in astrocytes, microglia, and macrophages, where it is a major source of reactive oxygen species (ROS) generation [4].

Snake venom (SV) from Vipera lebetina turanica has a group of basic peptides, and V activator (composed of 236 amino acids with 6 disulfide bonds formed by 12 cysteine) [5]. Raquel et al [6] elucidated that SV induced caspase cascade system apoptosis, and it has been demonstrated that SV has induced apoptosis with anti-cancer properties against colorectal cancer [7], prostate cancer cells [8], lung cancer [9,10], breast cancer [11] and ovarian cancer [12]. However, the molecular mechanisms of the anti-inflammatory effects of SV induced cannabinoid-infarction have not been reported. Thus, in the present study, the neuroprotective effects of SV were investigated using an animal model of human ischemic stroke [middle cerebral artery occlusion (MCAO) technique], to analyze anti-oxidative and anti-inflammatory, morphological and neurobehavioral profile changes in mice.

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Materials and Methods

Materials

The SV of Vipera lebetina turanica was purchased from Sigma (Saint Louis, MO). 7-week-old male ICR mice were purchased from Samtako BioKorea Co. (Osan, Korea).

Cell culture

Microglial BV-2 cells were (American Type Culture Collection, Rockville, MD) were incubated in 5% CO₂ at 37°C and grown to subconfluence. The medium used for routine culture was Dulbecco's modified Eagle's medium (DMEM, Invitrogen, Carlsbad, CA, USA), supplemented with 10% fetal bovine serum (FBS), and penicillin (100 U/mL).

Cells were incubated with SV at various concentrations (0.625, 1.25, or 2.5 μg/mL) and then stimulated with LPS 1 μg/mL. Various concentrations of SV dissolved in dimethyl sulfoxide (DMSO)-ethanol solvent (1:1 DMSO: ethanol, v/v) were added together with LPS. The final concentration of DMSO-ethanol solvent used was less than 0.05%. Cells were treated with 0.05% DMSO-ethanol solvent as vehicle control.

Microglial BV-2 cells were plated in 96-well plates, and subsequently treated with SV (0.625, 1.25, or 2.5 μg/mL) for 24 hours. After treatment, cell viability was measured using the MTT [3-(4, 5-Dimethylthiazol-2-yl)-2, 5-Diphenyltetrazolium Bromide] assay (Sigma Aldrich, St. Louis, MO, USA) according to the manufacturer's instructions. Briefly, MTT (5 mg/mL) was added to the culture plates and incubated at 37°C for 4 hours before 100 μg/mL dimethyl sulfoxide (DMSO) was added to each well. Finally, the absorbance of each well was read at a wavelength of 540 nm using a microplate reader.

Ischemia surgery

The acclimated 7-week-old ICR mice were narcotized with a gas mixture of 75% N₂O and 25% O₂. The middle cerebral artery was occluded for 1 hour using sutures (Doccol Corp, Sharon, MA, USA). The right common carotid artery and the right external carotid artery, were exposed through a middle neck incision. The right external carotid artery was dissected distally, ligated, and coagulated along with the terminal lingual and maxillary artery branches. A minimal incision was made in the right external carotid artery stump, at an angle of incidence, with iridectomy scissors. Following the incision, occlusion of the MCA was performed with sutures (0.22 mm). During the occlusion period and postoperative period (for 2 hours after occlusion), the animals were kept on thermostatically controlled warming plates in order to maintain body temperature at 37°C. Following the occlusion, clips were removed to restore blood recirculation. The animals surgically operated on without cerebral artery occlusion served as the Sham control. 1 hour after reperfusion, SV pharmacopuncture was applied intravenously to the caudal vein at a concentration of 50 ng/kg.

Morphometric determination of infarction

When the MCAO probe had been inserted for 24 and 72 hours, the brain tissue was stained with 2, 3, 5-triphenyltetrazolium chloride (TTC) to examine the ischemia infarction area. Briefly, the brain was removed and cut into 2 mm serial slices originating 1 mm from the frontal pole. The coronal slices were then immersed in a 2% phosphate-buffered solution (PBS) for 50 minutes at 37°C. After TTC staining, the slices were fixed in a 10% phosphate-buffered formalin and the area of infarction was then determined by an image analyzer using the Leica Qwin program (Leica Microsystems Imaging Solution Ltd., Cambridge, UK). The infarct area was fixed through an image program (Sion image, Scion Corporation, MD, USA), and then the infarct volume of the whole brain was calculated by the sum of all the slices (7 slices in 1 brain) of infarct area volume × thickness (2 mm). The relative infarction areas were indicated by the percentage of Sham control brain infarction areas.

Assessment of neurological deficit score

The neurological deficit scores were estimated 24 hours after the insertion of the MCAO probe as described by Longa et al. Scoring was based on the 5 types of disorder: (1) failure to extend the left forepaw, (2) decreased grip strength of forepaw, (3) circling left by pulling the tail, (4) spontaneous circling, (5) falling down. A rating of 1 point was given for each assessment, and then the scores were totaled. Neurological function assessment was performed by an investigator blinded to the experimental groups.

Nitrite quantification assay

Using ether anesthesia, the mice were sacrificed and cerebrospinal fluid was collected (taking care not to puncture the cervical cisterna membrane and contaminate the sample with blood). Nitrite in cerebrospinal fluid was measured as a final product of NO generation using a Griess reagent system at 540 nm (Promega, Madison, WI, USA).

ROS generation

Generation of ROS was assessed by 2, 7-dichlorofluorescein diacetate (DCFH-DA, Sigma Aldrich), an oxidation-sensitive fluorescent probe. Intracellular H₂O₂ or low-molecular-weight peroxides, oxidize DCFH-DA to DCF, a highly fluorescent compound. Briefly, tissue lysates were plated in black 96 well plate, and incubated with 10 μM DCFH-DA, at 37°C for 4 hours. The fluorescence intensity of DCF was measured in a microtiter plate reader at an excitation wavelength of 485 nm, and an emission wavelength of 538 nm.

Assay for glutathione peroxidase activity

Five half brains were used to detect glutathione peroxidase activity. Brain tissue was sonicated in PBS for 15 seconds on ice, and the homogenate was centrifuged at 1,000 g at 4°C for 5 minutes. The supernatant was used in the glutathione peroxidase assay. Briefly, the GSSG produced during the glutathione peroxidase enzyme reaction was immediately reduced by glutathione reductase, and reduced nicotinamide adenine dinucleotide phosphate (NADPH). Therefore, the rate of NADPH consumption (monitored as a decrease in absorbance at 340 nm) was proportional to formation of GSSG during the glutathione peroxidase reaction. The reaction buffer contained 20 mM potassium phosphate, (pH 7.0), 0.6 mM ethylene diamine tetraacetic acid, 0.15 mM NADPH, 4 U of glutathione reductase, 2 mM GSH, 1 mM sodium azide, and 0.1 mM H₂O₂. This assay was performed at 25°C. 1 U of glutathione peroxidase activity was defined as 1 µM NADPH consumed per minute.

Real-time qPCR

Total RNA was extracted using Trizol reagent (Sangon, Shanghai,
China) from homogenized tissues or cells. Reverta Ace qPCR RT Kit (Toyobo, Osaka, Japan) was used to generate cDNA from mRNA, and SYBR Premix Ex Taq (Takara, Kyoto, Japan) was used for real-time qPCR with the ABI 7500 real-time qPCR system, following the manufacturer’s instructions. The primers used were mouse IL-1β, TNF-α.

Cell viability

The cell suspension was counted using a hemocytometer and 180 µL of the cell suspension was added at the desired concentration, to each well of 96 cell plate. The specimen was dissolved in PBS and 20 µL of each specimen was added to each well and incubated with 100 µL of MTT solution. After 4 hours of incubation, the MTT dilution was carefully removed, and 100 µL of DMSO was added to each well and shaken for 15-20 minutes. The ELISA reader measured absorbency at a wavelength of 540 nm. This absorbance represented the amount of MTT reduced by the cell, and was proportional to the number of surviving cells present in each well.

Western Blot analysis

The brain tissues were homogenized with lysis buffer (PROPREP; nNtRON, Sungnam, Korea; n = 8 mice per group), and centrifuged at 2,500 g, for 15 minutes, at 4°C. Equal amounts of total protein (40 µg) isolated from brain tissue were resolved on 8% or 10% sodium dodecyl sulfate polyacrylamide gels, and then transferred to nitrocellulose membranes (Hybond ECL; Amersham Pharmacia Biotech, Piscataway, NJ, USA). Membranes were incubated at room temperature for 2 hours with the following specific antibodies: anti-COX-2, anti-1KB, anti-p-1KB (Cell Signaling Mol Neurobiol Technology, Inc., Beverly, MA, USA), anti-iNOS (1:1,000 Novus Biologicals, Inc., Littleton, CO), Mouse monoclonal antibodies directed against JNK, phosphor-JNK, p38, phosphor-p38, ERK, phosphor-ERK (1:1,000) (Cell signaling Technology, Beverly, MA, USA) and anti-β-actin (1:2,500; Sigma, St Louis, MO, USA). Blots were then incubated at room temperature for 2 hours with the corresponding peroxidase-conjugated anti-mouse or anti-rabbit antibodies (1:2,000; Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA). Immunoreactive proteins were detected using an enhanced chemiluminescence Western blotting detection system. The relative density of the protein bands was scanned using My Image (SLB, Seoul, Korea) and quantified by Lab Works 4.0 (UVP, Upland, CA, USA) for density.

Microglial BV-2 cells (American Type Culture Collection (Rockville, MD, USA) were grown to subconfluence in an incubator at 5% CO₂, at 37°C. The medium used was DMEM, supplemented with 10% FBS, and penicillin (100 U/mL). Equal amounts of total protein (40 µg) isolated from Microglial BV-2 cells were resolved on 8% or 10% sodium dodecyl sulfate polyacrylamide gels and then transferred to nitrocellulose membranes (Hybond ECL; Amersham Pharmacia Biotech, Piscataway, NJ, USA). Membranes were incubated at room temperature for 2 hours with the following specific antibodies: anti-COX-2 (Cell Signaling Mol Neurobiol Technology, Inc., Beverly, MA, USA), anti-iNOS (1:1,000 Novus Biologicals, Inc., Littleton, CO, USA), Mouse monoclonal antibodies directed against JNK, phosphor-JNK, p38, phosphor-p38, ERK, phosphor-ERK (1:1,000) (Cell signaling Technology, Beverly, MA, USA) and anti-β-actin (1:2,500; Sigma, St Louis, MO, USA). Blots were then incubated at room temperature for 2 hours with the corresponding peroxidase-conjugated anti-mouse or anti-rabbit antibodies (1:2,000; Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA). Immunoreactive proteins were detected using an enhanced chemiluminescence Western blotting detection system. The relative density of the protein bands was scanned using My Image (SLB, Seoul, Korea) and quantified by Lab Works 4.0 (UVP, Upland, CA, USA).

Statistical analysis

The data were analyzed using the GraphPad Prism 4 software (GraphPad Software, La Jolla, CA, USA). Data were presented as mean ± SD. Differences in the data were assessed by a 1-way analysis of variance and when p = 0.05, the differences were assessed by Dunnett’s test. A value of p < 0.05 was considered to be statistically significant.

Results

The numbers of mice in the MCAO experimental group were reduced in this study because the numbers in the Sham operated group were reduced.

Effect of SV on cerebral infarction induced by MCAO

Cerebral infarction was observed in the brain of mice after MCAO, as observed by TTC staining of brain sections (Fig. 1A). MCAO. The MCAO+SV 50 ng/kg (SV 50 ng/kg treatment group) showed that treatment with SV reduced the level of infarction and the infarct area compared with the MCAO group (Figs. 1A and 1B), whereas the Sham group had no infarction (Fig. 1A).

Effect of SV on the levels of neurological deficit score

On the day of implementation, neither the experimental nor the control group showed significant changes in neurological deficit score, but after 24 hours, the SV treatment group had a neurological deficit score significantly lower than on the implementation day (p < 0.05; Fig. 2).

Effect of SV on the levels of NO and ROS

The SV (50 ng/kg) treatment group showed no significant difference in the concentration of nitric oxide compared with the
**Effect of SV on NO and ROS**

Fig. 3. Effect of SV on the levels of NO and ROS. (A) After 72 hours from MCAO probe insertion, infarction volume of brain tissue treated with SV I.V. (50 ng/kg) was made by mice sacrificed. After that, measuring NO and ROS, it indicates statistically significant differences from SV treated group. (B) After treatment of SV (50 ng/kg), ROS were decreased, but NO was not significantly different from MCAO group. MCAO, middle cerebral artery occlusion; SV, snake venom; TTC, triphenyltetrazolium chloride; NO, nitric oxide; ROS, reactive oxygen species.

The ratio of GSH/GSSG concentration indicated that treatment with SV (50 ng/kg) statistically significantly reduced the GSH/GSSG ratio compared with the MCAO group (Fig. 4; p < 0.05). When GSH/GSSG ratio decreased, cells oxidize. The GSH/GSSG ratio in the MCAO group decreased in contrast to the Sham group, and SV treatment group (Fig. 4).

**Effect of SV on the levels of IL-1β and TNF-α using qPCR**

Fig. 5. Effect of SV on the levels IL-1β and TNF-α using qPCR. (A) MCAO group were treated with SV I.V. (50 ng/kg). After treatment, measuring IL-1β or TNF-α, it indicates statistically significant differences from SV treated group. (B) After treatment of SV (50 ng/kg), IL-1β or TNF-α were significant decreased. IL-1β or TNF-α convergence has been reduced by the group that has handled SV in MCAO. **p < 0.01, indicates statistically significant differences from SV treatment group. ***p < 0.001, indicates statistically significant differences from SV treatment group. MCAO, middle cerebral artery occlusion; SV, snake venom; qPCR, quantitative polymerase chain reaction.

IL-1β and TNF-α also activates the transcription of the iNOS gene through NF-kB activation. Fig. 5 shows that IL-1β and TNF-α concentrations were decreased significantly by treatment with SV compared with the MCAO group (p < 0.01; Figs. 5A and 5B).

**Effect of SV on the expression of enzymes caused by MCAO**

The relationship between the expression of related regulatory enzymes in the MCAO group treated with SV (50 ng/kg) was investigated using Western blots. The expression of iNOS and COX-2 was reduced in the SV group compared with the MCAO group alone. Expression of β-actin protein was used as an internal control (Fig. 6).

The expression of inflammatory enzymes was also examined between groups. Treatment with SV reduced the expression of p-IκB-α, P38, p-JNK, p-ERK compared with the MCAO group.
Effects of SV on the expression of enzymes caused by MCAO. 

The expression of protein enzymes was determined using Western blot analysis. The MCAO group were treated with SV I.V. (50 ng/kg) for 24 hr. Equal amounts of total proteins (50 ng/kg) were subjected to 12% or 8% SDS-PAGE. Expression of iNOS, COX-2 and β-actin were detected by Western blotting using specific antibodies. β-actin protein here was used as an internal control. Each band is representative for three experiments.

MCAO, middle cerebral artery occlusion; SV, snake venom; iNOS, inducible nitric oxide synthase; COX-2, cyclooxygenase-2.

Expression of iNOS was increased by the treatment with SV in a concentration dependent manner (Fig. 7).

In addition, expression of apoptosis regulatory proteins was determined using Western blot analysis. MCAO were treated with different concentrations of SV. Each band is representative for three groups.

MCAO, middle cerebral artery occlusion; SV, snake venom; p-IkB-α, phospho-inhibitor kappa beta-alpha; p-JNK, phospho-Jun N-terminal kinase; p-ERK, phospho-extracellular signal-regulated kinase.

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Effect of SV on LPS-induced inflammation-state BV2 cell growth

The inhibitory effect of SV in the presence of LPS (1 µg/mL) on cell growth of human keratinocyte BV2 cells was investigated. Cell viability by MTT assay and cell count was analyzed. The cells were treated with a concentration range of SV (0, 0.313, 0.625, 1.25, 2.5, 5, 10, 20, 40 µg/mL) for 24 hours in the presence of LPS. BV2 cells survived up to a concentration of 20 µg/mL of SV (Fig. 8).

Expression of enzymes induced by MCAO according to the different concentrations of SV

The enzymes produced during inflammation in the brain cells of the MCAO mouse model during treatment with different concentration of intravenous SV (50 ng/kg) was investigated. The cells were treated with 4 concentrations of SV (0, 0.625, 1.25, 2.5 µg/mL) for 24 hours with or without the presence of LPS. Cox-2
was expressed at the highest level without external stimulation with LPS or SV. Treatment with 2.5 μg/mL SV reduced the expression of p-JNK and p-IκB-α compared with 1.25 μg/mL and 0.625 μg/mL SV, and no SV (Fig. 9).

Discussion

Cerebral ischemia reduces cerebral blood flow and oxygen supply, and leads to neuronal cell death. This process is caused by energy depletion and inflammatory reactions. When cerebral ischemia occurs, neurons depolarize, synaptic glutamate is released, and glutamate accumulates extracellularly. Furthermore, extracellular glutamate accumulation is accelerated by the reversal of active glutamate transport, due to reduced ATP levels. This extracellular glutamate released by Ca\(^{2+}\)-dependent exocytosis induces excessive intracellular Ca\(^{2+}\) influx largely through potentiated NMDA-receptor channels, which leads to free radical production via the activation of Ca\(^{2+}\)-dependent protease, lipase, and various other modulators. Eventually, these free radicals destroy the cellular structure and induce neuronal death. In addition, decreases in intracellular polyamines, which are known to scavenge free radicals [13] and an increase in hydroxyl radicals [14], are also evident after focal cerebral ischemia. Furthermore, NO, which is known to regulate metabolism in the brain and blood flow, and participate in NMDA-receptor-mediated neural toxicity, has been reported to have a significant influence on cerebral ischemia [15]. The inflammatory reactions of ischemia are preceded by an accumulation of neutrophils, an increase in cytokines, and an increase in white blood cell adhesion molecules, and astrocyte numbers. During the early stage of the post-ischemic inflammation reaction, granulocytes, such as, neutrophils and eosinophils, are recruited from blood, whereas during the late stage, T cells and macrophages infiltrate ischemic areas. Consequently, neuronal cell damage occurs and is irreversible, even if blood circulation has been fully restored. It has been reported that microglia inflammation is more prominent in focal cerebral ischemia in ICR rats than in normal rats [16].

SV is very dangerous, but many researchers report that natural SV is a useful biological resource, containing several pharmacologically active components that could be of potential therapeutic value [17-19], and Song et al demonstrated that SV from Vipera lebetina turanica inhibited NF-κB activation, and growth of cancers such as neuroblastoma and prostate cancer, through its interaction with signal molecules in the NF-κB signal pathway [7,8].

TLR4 detects LPS coupled with CD14, and signals are generated inside the cell, iκB which is phosphoric acidized by iκK, become degraded, the free NF-κB enters the nucleus and transfers the iNOS protein, producing excessive NO, and releasing inflammatory cytokines into the blood. In addition, COX-2 protein will be transferred to help in the composition of prostaglandins, which will lead to inflammation [20]. SV maintains iκB levels which prevents NF-κB activation and the spread of iNOS and COX-2, ultimately eliminating the spread of inflammation. During inflammation, tissues are oxidized, as seen in the ROS and GSH/ GSSG assays in this study. SV treatment prevented oxidation and, as seen in qPCR, reduced the level of inflammatory cytokines. This can be observed in TTC staining, and it can be determined that SV may be involved in the recovery of damaged tissue. In this study, a moderate concentration of SV may be useful for recovering damaged tissue (2.5 to 20 μg/mL as observed in Western blot and cell viability assays).

It is well known that activation of astrocytes is a result of brain injury. Thus, the increased expression of astrocyte cytoskeletal protein during neural tissue injury could be a quantitative marker [21]. As predicted, SV decreased the number of activated astrocytes in the subventricular zone and striatum, in parallel with decreased expression of in the infarction area. A neuroprotective effect due to SV, is inferred from the antioxidant and anti-inflammatory activities of pyrogallol, a major component of SV, which strongly scavenges free radicals, and inhibits inflammatory mediators [22,23].

Exerted neuroprotective effects in a MCAO model of stroke are caused by suppressing astrocyte activation and inhibiting inflammatory and oxidative indices. Although the active ingredients in SV remain unidentified, SV improved neurobehavioral functions via neuroprotective activity. Therefore, it was concluded that SV may be a good neuroprotective agent, suggesting it may be advantageous for pharmaceutical applications for ischemic stroke.

In conclusion, in the present study, SV was found to repress cerebral ischemia regardless of preexisting conditions or complications, and all animals treated with SV showed a reduction in cerebral ischemia. The main treatment target after cerebral ischemia, is not necrosis of the core, but rather apoptosis of the penumbral region. The findings in this study indicate that SV effectively ameliorated neuron damage in the penumbral region, and markedly reduced ischemic damage in ICR mice. In view of the reduced levels of iNOS and COX-2, SV might attenuate infarct areas via an anti-inflammatory cytokine related effect through the reduction in NF-κB activity.

Conflicts of Interest

The authors have no conflicts of interest to declare.

References


