Neuroprotective Effect of Visnagin on Kainic Acid-induced Neuronal Cell Death in the Mice Hippocampus

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Visnagin (4-methoxy-7-methyl-5H-furo[3,2-g][1]-benzopyran-5-one), which is an active principle extracted from the fruits of Ammi visnaga, has been used as a treatment for low blood-pressure and blocked blood vessel contraction by inhibition of calcium influx into blood cells. However, the neuroprotective effect of visnagin was not clearly known until now. Thus, we investigated whether visnagin has a neuroprotective effect against kainic acid (KA)-induced neuronal cell death. In the cresyl violet staining, pre-treatment or post-treatment visnagin (100 mg/kg, p.o. or i.p.) showed a neuroprotective effect on KA (0.1 μg) toxicity. KA-induced gliosis and proinflammatory marker (IL-1β, TNF-α, IL-6, and COX-2) inductions were also suppressed by visnagin administration. These results suggest that visnagin has a neuroprotective effect in terms of suppressing KA-induced pathogenesis in the brain, and that these neuroprotective effects are associated with its anti-inflammatory effects.

Key Words: Neuroprotection, Kainic acid, Hippocampus, Visnagin, Cytokines

INTRODUCTION

Kainic acid (KA), which is an analog of the excitatory amino acid L-glutamate, elicits neuronal cell death followed by severe status epilepticus in the pyramidal layer of the hippocampal CA3 region [1] when KA is administered intracerebroventricularly (i.c.v.). It may be due to excessive activation of neurons by excitatory neurotransmitters (e.g. glutamate), which are massively released as a consequence of energy depletion and which result in excitotoxic neuron death [2,3]. Recent studies have demonstrated that the KA-induced neuronal death is associated with the activation of microglia and astrocytes in the hippocampus, and that these processes are induced by enhanced reactive oxygen species (ROS) production and cytokine expressions [4,5]. The microglia also can be detected using the complement receptor type 3 (OX-42) IR. The complement receptor type 3 is important in the adherence of neutrophils and monocytes to stimulated endothelium, and also in the phagocytosis of complement coated particles that raises the last step of the microglial activation [6]. Expressions, hypertrophy and proliferation of Glial fibrillary acidic protein (GFAP) specifically is found in astroglia, a cell type which is highly responsive to neurologic insults [7]. After brain injury, astrocytes undergo a number of cellular syntheses and release of a variety of growth factors and immunomodulatory cytokines [7]. In addition, recent studies have demonstrated that inflammatory and apoptotic processes contribute to the later stages of the damage induced by various brain injuries, and that these detrimentally affect neurologic outcome [5,8]. Among them, it has been reported well that pro-inflammatory mediators such as IL-1β, TNF-α, IL-6, and COX-2 can lead to deteriorative effect in the brain, and KA can trigger an aberrant inflammatory cytokine response by microglial cells and accelerated disease progression [4].

Visnagin (4-methoxy-7-methyl-5H-furo[3,2-g][1]-benzopyran-5-one) is an active principle extracted from the fruits of Ammi visnaga [9]. The fruit or its isolated active components have been used for the treatment of angina pectoris due to their peripheral and coronary vasodilator activity [10,11]. In isolated aorta, visnagin, and other related active principles present in these fruits such as visnadin and khellin inhibited vascular smooth muscle contractility, probably by acting at multiple sites to decrease the availability of Ca2+ required for activation [12-14]. Several reports have demonstrated that visnagin descents blood pressure and blocks blood vessel contraction as inhibiting calcium influx into cell [15]. However, the neuroprotective effect of visnagin on kainic acid (KA)-induced neuronal death has not been clearly demonstrated yet.

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ABBREVIATIONS: KA, kainic acid; GFAP, glial fibrillary acidic protein; IR, immunoreactivity; IL-1β, interleukin-1beta; TNF-α, tumor necrosis factor-alpha; IL-6, interleukin-6; COX-2, cyclooxygenase-2.
been demonstrated until now although visnagin has been studied for therapeutic use for over 10 years.

In the present study, we explored the neuroprotective effects of visnagin in KA-induced neuronal cell death model. The peroral (p.o.) or intraperitoneal (i.p.) administration of visnagin remarkably suppresses hippocampal cell death, indicating that visnagin has a neuroprotective effect against KA-induced neuronal cell death.

**METHODS**

These experiments were approved by the Hallym University Animal Care and Use Committee. All procedures were conducted in accordance with the 'Guide for Care and Use of Laboratory Animals' published by the National Institutes of Health. Male ICR mice (MJ Co., Seoul, Korea) weighing 25-28 g were used for all the experiments.

**Experimental animals**

Male ICR mice (MJ Co., Seoul, Korea) weighing 25-28 g were used for all the experiments. Animals were housed 5 per cage in a room maintained at 22±0.5°C with an alternating 12 hr light-dark cycle. Food and water were available *ad libitum*. The animals were allowed to adapt to the laboratory for at least 2 hr before testing and were only used once. To reduce variation, all experiments were performed during the light phase of the cycle (10:00~17:00).

**Drug treatment and i.c.v. kainic acid injection**

Visnagin was purchased from Acros organics. Visnagin was prepared following steps: (A) 1 g of decursinol was dissolved in 0.5 ml of ethanol plus 0.5 ml of polyethylene glycol 400 (B) Separately; 100 mg of sodium carboxymethyl-cellulose was dissolved in 9 ml of distilled water. (C) Finally, Solution (A) and Solution (B) were vigorously mixed. These solutions excluding visnagin were used as vehicle control. The KA (Sigma, USA) was dissolved in a phosphate buffer solution. The i.c.v. administrations of KA were performed following the procedure established by Laursen and Belknap [16]. Briefly, each mouse was injected at bregma with a 50 μl Hamilton microsyringe fitted with a 26-gauge needle that was inserted to a depth of 2.4 mm.

**Cresyl violet staining method and histological analysis**

Animals were sacrificed for the brain sample by perfusion at 1 day after KA administration. All perfusion procedures were worked in the fume hood. For perfusion, all mice were first deeply anesthetized with sodium pentobarbital (100 mg/kg, i.p., Hanlim Pharm., Korea) and perfused intracardially with physiological saline followed with ice-cold phosphate-buffered 4% paraformaldehyde (pH 7.4). Whole brain was removed from the skull and postfixed in the same fixative for 4 hrs at 4°C. Then the brains were cryoprotected in 30% sucrose for 24 hrs at 4°C and sectioned coronally (45 μm) on a freezing microtome and collected in cryoprotectant for storage at −20°C until processed. Prepared sections were rinsed 3×10 min in PBS to remove cryoprotectant. Sections were mounted on microscope slides (Fisher, USA) and dried on air. The slides were soaked in cresyl violet working solution (0.02% buffer solution; 0.2% sodium acetate, 0.3% acetic acid) for 30 min. Then the sections were dehydrated through alcohol and xylene and covered slipped using Permount (Fisher, USA).

Histological analysis method in pyramidal layer of hippocampal CA3 region was performed following under procedures [17]. The number of cresyl violet-positive neurons was counted by two blinded observers at the same time using an image analyzing system equipped with a computer-based CCD camera (Olympus AX70, USA). The number of cresyl violet-positive neurons in the CA3 region of the hippocampus was counted in 3 sections in reference to the mouse atlas [18] for each animal [19]. Starting from the first section (interaural 2.10 mm, bregma −1.70 mm), counts were taken from at least three coronal sections at 0.135 mm increments. Thus, we could always perform neuronal counting of the same brain region and minimize any counting bias. The number of cresyl violet-positive neurons was compared to that of the control group of the same brain area from all animals. All experiments were conducted independently twice. The neuronal counting of the same group was combined for final analysis.

**Isolation of total RNA**

The entire process for isolation of total RNA was conducted three times independently. Finally, the animal number used for each group was nine. Three animals of each group were dissected for real time PCR analysis. Total cellular RNA was extracted from dissected hippocampus tissue using a rapid guanidine thiocyanate-watersaturated phenol/chloroform extraction procedure and subsequent precipitation with acidified sodium acetate [20]. Total cellular RNA in the aqueous phase was precipitated with ice-cold isopropl alcohol. Isolated RNA samples were subjected to spectrophotometric analysis at 260 nm and 280 nm. The separated organic layer was extracted twice with an equal volume of sterilized (Millipore, USA) water and proteins were precipitated by adding two volumes of absolute ethanol to the water-extracted organic phase. The dried pellets were dissolved in a denaturing buffer (6 M guanidinium chloride, 20 mM Tris-HCl [pH 8.0], and 1 mM EDTA).

**Real-time PCR analysis**

Expression of TNF-α, IL-1β, IL-6, COX-2 and GAPDH mRNA was evaluated by real-time PCR using QuantiTect SYBR® Green PCR Kit (Qiagen, Germany). All PCRs were performed in total volume of 20 μl using the QuantiTect SYBR® Green PCR Kit (Qiagen, Germany). Each reaction contained 1.5 μl of cDNA, 6.5 μl RNase-Free Water, each 1 μl of sense and antisense primer (20 μM) and 10 μl of 2X SYBR® Green PCR Master Mix (containing QuantiTect SYBR Green PCR buffer, dNTPs, SYBR Green I dye, ROX dye, and HotStarTag DNA polymerase). After an initial denaturation step at 95°C for 30 s, temperature cycling with a total of 40 cycles was initiated. Each cycle consisted of a denaturation phase at 95°C for 30 s, an annealing phase at 60°C for 30 s and an elongation phase at 72°C for 30 s. Amplification was followed by melting curve analysis to verify the correctness of the amplification. A negative control with water instead of cDNA was run within every PCR to assess specificity of the reaction. To verify the accuracy of the amplification, PCR products were further analyzed on ethidium bromide-stained 2% agarose gel. For data analysis, Rotor-Gene 6000 Series Software 1.7 (Build 87) was used. Results are given as a ratio of the amount of TNF-α,
Neuroprotective Effect of Visnagin

Fig. 1. The effect of visnagin administered orally or intraperitoneally on KA-induced neuronal death in the hippocampus. The location of the hippocampal CA3 region performing cresyl violet-positive neuronal count is indicated (A) referring Franklin [18]. I.c.v. kainic acid injection induced neuronal cell death in the pyramidal cells in the CA3 region of hippocampus (B). Mice were administered visnagin orally (C) or intraperitoneally (D) 20 min (− 20) prior to KA (0.1 μg/5 μl) injection or 0, 20 (+20), 40 (+40), 60 (+60) min after KA treatment (0.1 μg/5 μl). And then, the cresyl violet staining was performed at 1 day after KA. The vertical bars indicate the standard error of mean. *p < 0.05, **p < 0.01, ***p < 0.001 (KA 0.1 vs other groups). The mice number of each group was 10.
duced neuronal death, we administered visnagin orally (p.o.) or intraperitoneally (i.p.) 20 min (−20 min) prior to KA (0.1 μg/5 μl, i.c.v.) injection or 0, 20 (+20), 40 (+40), 60 (+60) min after KA injection (0.1 μg/5 μl, i.c.v.). As shown in Fig. 1A, KA induced neuronal cell death in the pyramidal layer of the hippocampal CA3 region. When visnagin was treated before or after KA injection orally or intraperitoneally, we observed that visnagin had neuroprotective effect on KA-induced neuronal cell death. However, delayed visnagin administration (+40 and +60 min) in both p.o. (Fig. 1B) and i.p. (Fig. 1C) did not show any neuroprotective effect on KA-induced neuronal death. The visnagin administered p.o. or i.p. at −20, 0, +20 min had similar neuroprotective effect on KA-induced neuronal death in the hippocampus.

The course alteration of proinflammatory cytokines mRNA on KA injection in the hippocampus

Previous studies suggest that inflammatory and apoptotic processes contribute to the later stages of the damage induced by brain injuries, and that these detrimentally affect neurologic outcome [5,21,22]. Therefore, we investigate the course alteration of proinflammatory cytokines mRNA on KA injection in the hippocampus. The mRNA levels of various proinflammatory markers such as IL-1β, TNF-α, IL-6, and COX-2 in the hippocampus were examined at 1, 3, 6, 12, 24 hrs after KA administration (0.1 μg/5 μl, i.c.v.).

As shown in Fig. 2, intracerebroventricular injection with KA increased TNF-α and COX-2 mRNA levels, and the levels showed peak at 3 hrs after KA treatment. On the other hand, IL-1β mRNA level reached maximum level at 1hr after KA treatment, and IL-6 mRNA level showed peak level at 3 and 6 hrs after KA treatment. The increased cytokines mRNA level decreased gradually after peak level.

The effect of visnagin on pro-inflammatory cytokines increased by KA in the hippocampus

To investigate whether the neuroprotective effect of visnagin is accompanied by suppressions of proinflammatory markers such as IL-1β, TNF-α, IL-6, and COX-2 in the hippocampus, we observed an effect of visnagin on pro-inflammatory cytokines increased by KA in the hippocampus. The time point when pro-inflammatory cytokines peaked was referred by Fig 2 result. The induction of IL-1β, TNF-α, IL-6, and COX-2 mRNA hippocampus were inhibited by visnagin pretreatment (Fig. 3). However, visnagin itself did not affect pro-inflammatory cytokines expression.

The effect of visnagin on the GFAP, OX-42 expression induced by KA in the hippocampal CA3 region

To investigate whether visnagin affects GFAP or OX-42 expression induced by KA, the activations of microglia and astrocytes were analyzed using anti-OX-42 (microglia marker) and anti-GFAP (astrocyte marker) antibodies, respectively (Fig. 4). In PBS treated control mice, both microglia and astrocytes having ramified morphology were barely detected. However, the number of anti-OX-42 cells and anti-GFAP cells was remarkably elevated at 1 day after i.c.v. KA treatment in the hippocampal CA3 region. In addition, visnagin administered orally at −20 min attenuated OX-42 and GFAP expression induced by KA.

DISCUSSION

There is an accumulating body of evidence which suggests that inflammation contributes to brain damage occur-
ring after acute injury, and that it detrimentally affects neurological outcome [5,8]. In addition, under inflammatory conditions, free oxygen radicals, nitric oxide, and inflammatory cytokines produced by activated microglial cells seem to cause neuronal damage. Because the hippocampus is densely populated with microglia and is one of the most sensitive and malleable regions of the brain [23], it is speculated that excessive production of inflammatory cytokines in the hippocampus would be associated with KA toxicity. In the present study, we observed that pre- or post-visnagin administered orally (p.o.) or intraperitoneally (i.p.) inhibited KA-induced neuronal cell death in the hippocampal CA3 region. Visnagin not only inhibited microglial and astroglial activation but also attenuated the inflammatory marker expressions concomitantly, suggesting that visnagin exerts its neuroprotective effects via an anti-inflammatory mechanism in KA model.

Visnagin has been shown to relax KCl- and noradrenaline-induced contractions in guinea-pig aortic strips to a similar extent [13] and this vasorelaxant effect was explained by the inhibition of Ca\(^{2+}\) entry into vascular smooth muscle cells. During excitotoxicity, glutamate concentrations increase in the synapse leads to intracellular Ca\(^{2+}\) concentration increase, activating a cell death pathway [2,3,24]. Thus, it is speculated that visnagin may have a neuroprotective effect against KA toxicity by inhibiting Ca\(^{2+}\) influx. Although inhibition of visnagin on Ca\(^{2+}\) entry into hippocampal neuron was not confirmed in the present study, it may be one mechanism on the neuroprotective effect of visnagin. In regard to its anti-inflammatory effect, we also observed whether visnagin attenuated inflammatory markers against KA toxicity in the hippocampus. It appears that khellin extracts have some antimicrobial activity; this might be attributable to both the khellin and visnagin constituents, which both seem to have antifungal, antibacterial, and antiviral activity [25]. In addition, several reports have demonstrated that drugs or compounds having an anti-inflammatory effect can show a neuroprotective effect [4,26,27]. Thus, we examined the effect of visnagin on proinflammatory mediators expression induced by KA.

Recently, it has been demonstrated that proinflammatory cytokines have established potent pro-excitatory actions [28-31]. Collectively these reports show that proinflammatory cytokines simultaneously facilitate excitatory glutamatergic pathways while concurrently reducing inhibitory GABAergic transmission. Thus it is speculated that the combinatorial synergic effect of these cytokines increased by KA and known facilitatory action of KA on excitatory neurotransmission may lead to neuronal cell death in the hippocampus. Thus, the inhibitory effect of visnagin on cytokines expression showing synergic effect on neuronal cell death by KA may explain the neuroprotective effect of visnagin. It has been reported that COX-2 inhibitors may protect the brain against neurodegenerative diseases [32]. In addition, COX-2 specific inhibitors have a neuroprotective effect in models of focal [33] and global brain ischemia [34]. Furthermore, reactive oxygen species are generated by COX-2 activity. Oxidative stress has been demanded to induce neurodegeneration in a variety of disease states [35]. Prostaglandins, which are the product of cyclooxygenase metabolism, can produce injury by inflammatory and vascular mechanisms, as well as directly lead to apoptosis in some cell types [36]. Furthermore, it has been reported that antioxidative effects were also observed in the case of visnagin [37]. Thus, it is speculated that COX-2 inhibition and the antioxidative effect of visnagin may be another mechanism explaining the neuroprotective effect on KA toxicity.

Several earlier reports showed microglia expressing many proinflammatory cytokines after KA treatment [28-31]. Collectively these reports show that proinflammatory cytokines simultaneously facilitate excitatory glutamatergic pathways while concurrently reducing inhibitory GABAergic transmission. Thus it is speculated that the combinatorial synergic effect of these cytokines increased by KA and known facilitatory action of KA on excitatory neurotransmission may lead to neuronal cell death in the hippocampus. Thus, the inhibitory effect of visnagin on cytokines expression showing synergic effect on neuronal cell death by KA may explain the neuroprotective effect of visnagin. It has been reported that COX-2 inhibitors may protect the brain against neurodegenerative diseases [32]. In addition, COX-2 specific inhibitors have a neuroprotective effect in models of focal [33] and global brain ischemia [34]. Furthermore, reactive oxygen species are generated by COX-2 activity. Oxidative stress has been demanded to induce neurodegeneration in a variety of disease states [35]. Prostaglandins, which are the product of cyclooxygenase metabolism, can produce injury by inflammatory and vascular mechanisms, as well as directly lead to apoptosis in some cell types [36]. Furthermore, it has been reported that antioxidative effects were also observed in the case of visnagin [37]. Thus, it is speculated that COX-2 inhibition and the antioxidative effect of visnagin may be another mechanism explaining the neuroprotective effect on KA toxicity.

Several earlier reports showed microglia expressing many inflammatory mediators after inflammatory injury [38,39]. Recently, it has been reported well that activated microglia is a significant source of redundant extracellular glutamate that induces excitotoxic neuronal death [40]. In addition,
there is a report that excitatory amino acids released by microglia are suggested to compose the major determinant of neurotoxicity rather than reactive oxygen intermediates and cytokines [41]. Furthermore, Liang et al. [42] have demonstrated that astrocytes prevented excito-neurotoxicity by the reduction of exogenous glutamate whereas microglia did not, and conversely, activated microglia released an excess of glutamate that induced excitotoxic neuronal death. Taken together, it is speculated that activated glial cells induced by KA may be involved in increasing extracellular glutamate concentration finally as regulating glutamate release each other, or induce synergic effect on KA toxicity. Although we can not confirm that visnagin plays important roles as glutamate scavenger or glutamate receptor blocker, it seems that visnagin contributes to partly attenuate glutamate release as inhibiting gliosis.

We also observed that visnagin co-treatment (at 0 hr, p.o.) with KA injection produced a similar protective effect with Pre- or post-treatment of visnagin in the CA3 region. This result suggests that in addition to the anti-inflammatory effects, visnagin may also be effective during the acute damage process. First, direct inhibition of Ca\(^{2+}\) influx into neurons may be one possibility as mentioned above [13] although the underlying mechanism remains to be explored in detail. Second, the antioxydative effect of visnagin may be another candidate on co-treatment effect [37]. However, further study should be conducted to elucidate these hypotheses.

In conclusion, based on all this information, we speculate that the neuroprotective effects of visnagin in vivo are the results of multiple mechanisms, and that one of these may be associated with the suppression of inflammatory processes.

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