Neuroprotective Effect of Memantine on Hippocampal Neurons in Infantile Rat Hydrocephalus

Öz: İnfant sıçanda kaolin ile oluşturulan hidrosefalide memantin uygulanmasının hipokampal nöronlar üzerindeki koruyucu etkisi araştırılmıştır.

YÖNTEM ve GEREÇ: Üç haftalık Sprague-Dawley türü sıçanların cisterna magna sıvısına kaolin soluciónu eklenerek hidrosefalik model oluşturulmuştur. Deneçlerde memantin, dozuna göre, yada normal suya verilmiştir. Deneçlerin mrt ve histolojik incelemeleri yapılmış, nitrik oksit sentaz aktivitesi ve hippocampal CA1, CA2 ve CA3 bölgelerinde nöron sayları hesaplanmıştır.

BULGULAR: Memantin verilmesiyle hidrosefalik modelin hippokampal CA1, CA2 ve CA3 bölgelerinde nöron sayısında azalma ve nitrik oksit sentaz aktivitesinde azalma gözlemlenmiştir.

SONUÇ: Bu deneyde, memantin diğer antiedamik ve antiöksiden gibi koruyucu etkileri sergilediği gibi, hidrosefalik modelinde hipokampal nöronlarda koruyucu etkiye sahip olduğu gösterilmiştir.

ANAHTAR SÖZÇÜLERİ: Memantin, Antiedamik, Antiöksidant, Hippokampus, Sıçan
INTRODUCTION

In the pathogenesis of hydrocephalus, besides the primary mechanical injury associated with ventricular dilatation, the presence of a mechanism of a secondary injury resulting from destructive and ischemic alterations in periventricular area was demonstrated both in experimentally-induced hydrocephalic animal models (3, 11, 14) and in congenitally hydrocephalic rats (23). It has been observed that as long as the hydrocephalic process progresses, consequent alterations emerge preferably at those areas sensitive to ischemia. There are few studies comprehensively describing the injury mechanism in hippocampus during adult hydrocephalus (6, 8, 16, 25). Published data indicating the negative impacts of the hydrocephalic process on the hippocampus of the immature brain are even fewer in number (17). The hippocampus, especially its CA1 region, is known to be very susceptible to hypoxia and ischemia, and is a relatively important target for the secondary neuronal injury originated from hydrocephalic process. There are data regarding the neurodegenerative effect of hydrocephalic process especially on the pyramidal layer of CA1 subunit of hippocampus (15).

It is well known that secondary neuronal injury mechanisms can be impacted by pharmacological means in order to minimize secondary neuronal death following cerebral trauma and ischemia. The use of neuroprotective agents for preventing the secondary neuronal loss in the hydrocephalic process is a relatively a new issue and has not been studied widely. Regarding the neuroprotection in hydrocephalus, there are only a few reports published in the literature (7, 11, 14).

In recent years, memantine, a non-competitive NMDA receptor antagonist, has been demonstrated to have protective effects on neuronal death due to transient frontal brain ischemia (2), hypoxic cerebral ischemia (4), spinal cord ischemia (9), neonatal cerebral focal ischemia (24) and retinal ischemia (21) in several animal models. It has also been shown to significantly decrease hippocampal neuronal cellular death that emerged one week after cerebral trauma in a cortical impact injury model (22).

In this study we investigated the potential neuroprotective effect of systemic memantine administration on hippocampal neurons in a rat model of newly onset childhood hydrocephalus. We hypothesized that during hydrocephalic process, memantine would have a protective effect on secondary neuronal injury in hippocampal neurons through its mechanism of preventing ischemia induced injury and neurotoxicity.

MATERIAL and METHODS

1. Animals, hydrocephalus induction and verification

Thirty Sprague-Dawley rat pups at age 3 weeks with a mean weight of 50 ± 10 grams were used in the study. The rats were provided by the Center of Reproduction of Animals for Experimental Studies of the University Medical School and the experiment was approved by the local Ethics Committee with the code number HAEK 20/2. All efforts were made to minimize suffering and the number of animals used. All surgical procedures were conducted and animals were cared for according to the Declaration of Helsinki. Rats were divided into 3 groups. Group 2 and Group 3 consisted of eleven rats and Group 1 (control group) consisted of eight rats. Anesthesia was induced in the rats by intramuscular injection of ketamine (10 mg/kg). Each rat's neck was shaved and a median cranio-cervical incision was done. Under binocular loupe magnification superficial and deep cervical muscles were separated off the midline and the atlanto-occipital membrane was identified. The membrane was penetrated at a right angle by a sterile insulin injector of 27 G introduced into the cisterna magna. The cerebrospinal fluid outflow from the injection site was observed. Then 0.05 ml of sterile kaolin (Kaolin hydrated Aluminum Silicate, Sigma Chemical Co., St. Louis, MO, USA) suspension (250 mg/ml in 0.9% saline) was slowly injected into the cisterna magna of the rats in Group 2 and Group 3. The animals in Group 1 received a sham injection consisting of needle insertion only. Cervical muscles and skin were closed with separate sutures. Pups were housed in standard cages (four animals/cage) and provided with a normal 12-hour day/night lighting schedule with free access to water and pellet food. After 24 hours, a single daily dose of 20 mg/kg of memantine (Ebixa®, Lundbeck Inc.) solution prepared with 0.9% saline was injected into the peritoneum (Kaolin hydrated Aluminum Silicate, Sigma Chemical Co., St. Louis, MO, USA) suspension (250 mg/ml in 0.9% saline) was injected into the peritoneum of the rats in Group 3. The first and the second group were injected the same volume of 0.9 % saline solution. Injections were made once daily for two weeks. At the end of the two-week period, radiological verification of the hydrocephalic process was done. Magnetic resonance imaging (MRI) was performed with a 1.5 Tesla MR device (Philips Intera) via the use of C3 coil circular surface to obtain T2-weighted images of the brain in the coronal plane. The lateral ventricles and cerebrum were evaluated in the rostral cerebrum immediately anterior to the third ventricle to verify ventricular dilatation. Following MRI evaluation, the rats were decapitated and their brains were removed. All the brains were transected at the bulboemodular junction and then sectioned sagittally midline into two hemispheres. After labeling, all specimens were sent for histological and immunohistochemical analyses in a single-blinded fashion.

Histological analysis

Each hemisphere was sectioned coronally (10 μm thick at an interval of 320 μm) and the sections were stained with Hematoxylin-Eosin. Average neuronal counts in the CA1, CA2 and CA3 regions of hippocampal area were assessed under a light microscope (Olympus U-MDOB3). For each animal, average neuronal counts were obtained by counting four serial coronal sections at X400. Only complete neuronal cells with a clearly defined cell body and nucleus were counted as described in a previous study (22).

Immunohistochemical analysis

Tissue sections were deparaffinized and wet through graded alcohol. Endogenous peroxidase activity was blocked by incubation in 3% hydrogen peroxide for 20 min. After rinsing.
with distilled water, the sections were blocked with 1:10 normal goat serum to minimize nonspecific background staining. After protein blockage, the sections were incubated with u-NOS antibody for 60 minutes. The sections were then incubated in biotinylated goat anti-polyvalent for 20 minutes, and then in streptavidin peroxidase for 20 minutes. The slides were washed in running tap water and counterstained with HE for 20 seconds. Sections were washed again in running tap water and incubated with Lithium carbonate. They were then washed in water and covered.

For Nitric Oxide Synthase (NOS) immunostaining, Nitric Oxide Synthase–Universal (u-NOS) antibody (Lab Vision -RB-9261-R7) was used. Immunoreactivity of NOS in hippocampal regions was observed at 400x and was graded according to the five-stage staining score system as described elsewhere (13) as 0: No staining, 1: stain concentration under 25%, 2: stain concentration between 25%-50%, 3: stain concentration between 50%-75% 4: stain concentration over 75%.

2. Statistical analysis

Statistical analysis was performed by using the computer software program SPSS for Windows (SPSS Inc.). Neuron count values of the groups were analyzed by analysis of variance (ANOVA) with post hoc Tukey test calculations for intergroup comparisons. Statistical comparisons for NOS activity were made using Pearson’s chi-square test. All p-values less than 0.05 were considered to be statistically significant. The data of this study were expressed as a mean value ± standard error of mean (SEM).

RESULTS

Three rat pups (two in Group 2 and one in Group 3) that had complications during the hydrocephalic induction procedure and/or failed to develop hydrocephalus were excluded from the experimental protocol. The remaining 27 animals were classified as follows: Group 1 (non-hydrocephalic sham operated rats): n=8, Group 2 (non-treated hydrocephalic rats): n=9 and Group 3 (memantine treated hydrocephalic rats): n=10.

Radiological evaluation

In MRI examinations, T2-weighted images of the brain in the coronal plane were obtained. The lateral ventricles and cerebrum were evaluated in the rostral cerebrum immediately anterior to the third ventricle to verify ventricular dilatation. MRI studies revealed that one rat from Group 2 and one from the Group 3 failed to develop ventricular dilatation and those subjects were excluded from the study. Figure 1 shows an example of MR image displaying ventricular dilatation two weeks after the hydrocephalus induction. Figure 2 displays the MR image of a rat from Group 1 having normal ventricular structure.

Neuron counts

For each group, average number of spared neuronal cells with a clearly defined cell body and nucleus in CA1, CA2 and CA3 regions of hippocampus was determined. Table I summarizes the data and the results of the statistical analyses done. Briefly, two weeks after the induction, the hydrocephalic process seemed to significantly decrease the average number of spared neurons in all regions of the hippocampus both in Group 2 and in Group 3. Memantine treated rat pups showed significantly higher number of spared neuron counts in CA1 and CA2 regions compared with the non-treated subjects. Figure 3 shows the localization of the CA1, CA2 and CA3 regions of hippocampus in the rat pup. Figure 4 shows an example of histological specimen revealing spared and degenerated neuronal structures in CA1 region.

Nitric oxide synthase (NOS) immunoreactivity

In each group, immunoreactivity of NOS in the CA1, CA2 and CA3 regions of the hippocampus was scored according to the
five-stage grading system (13). Table II summarizes the data and the results of the statistical analyses performed. In short, our data showed that hydrocephalic process significantly increased the NOS immunoreactivity in CA1, CA2 and CA3 regions in Group 2 and in Group 3. The increase in NOS immunoreactivity was found to be significantly reduced in CA1 and CA2 regions of the rats which had received memantine treatment during the hydrocephalic process compared with the non-treated rats. Figure 5 (I, II, III) shows examples of photomicrographs displaying immunoeexpression of NOS for each group.

**DISCUSSION**

There are few studies comprehensively describing the injury mechanism in hippocampus during hydrocephalus (6, 8, 16, 17, 25). It has been observed that as long as the hydrocephalic process progresses, consequent alterations emerge preferably at those areas sensitive to ischemia. There are data regarding especially the effect on the pyramidal layer of the CA1 subunit of the hippocampus (15). Decreased neurofilament and synaptophysin immunoreactivity in the hippocampus has been accepted as an indicator of neuronal degeneration in experimentally induced hydrocephalus (20).

In the hydrocephalic process, the decline of synaptophysin immunoreactivity in the hippocampus was suggested to remind the phenomenon of ‘delayed neuronal death’, which is observed secondary to transient global ischemia (15, 19). In their study, Klinge et al. reported a global increase in NOS immunoreactivity in hippocampal neurons in kaolin-induced hydrocephalic adult rats two weeks after the induction. This increase was demonstrated to be in correlation with decreasing of cerebral blood flow at the early stage of hydrocephalus (16). They postulated that the global increase in hippocampal NOS staining of hydrocephalic subjects appeared to indicate a toxic neuronal response (16). Considering the important role of neuronal NOS enzyme in the acute excitotoxic injury of the central nervous system, an excitotoxic injury could be mentioned in the hydrocephalic process at hippocampus. In a previous study it has been well documented that neuronal NOS activation was secondary to an elevation of extracellular glutamate (10). The observations of Klinge et al. (16) seem to be in agreement with the findings of others (18) who found an increase in extracellular glutamate transporters in the short-term period after the induction of kaolin hydrocephalus. All these data show that the hydrocephalus process seem to affect hippocampal cells by the mechanism

**Table I:** Average Numbers of Spared Neuronal Cells with a Clearly Defined Cell Body and Nucleus in CA1, CA2 and CA3 Regions of Hippocampus for Each Group are Shown. The Data are Expressed as Mean Value ± Standard Error of Mean. The Data were Analyzed by Analysis of Variance (ANOVA) with Post Hoc Tukey Test Calculations for Intergroup Comparisons

<table>
<thead>
<tr>
<th></th>
<th>CA1</th>
<th>CA2</th>
<th>CA3</th>
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<tbody>
<tr>
<td>Group 1</td>
<td>105.75 ± 1.74</td>
<td>100.37 ± 2.13</td>
<td>94.50 ± 1.36</td>
</tr>
<tr>
<td>Group 2</td>
<td>79.13 ± 1.21</td>
<td>71.88 ± 5.02</td>
<td>69.61 ± 3.15</td>
</tr>
<tr>
<td>Group 3</td>
<td>88.43 ± 1.41</td>
<td>84.70 ± 2.12</td>
<td>73.10 ± 3.33</td>
</tr>
<tr>
<td>Group 1 – Group 2</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
</tr>
<tr>
<td>Group 2 – Group 3</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
<td>p &gt;0.05</td>
</tr>
<tr>
<td>Group 1 – Group 3</td>
<td>p &gt;0.05</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
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*All p-values less than 0.05 were considered to be statistically significant.

**Table II:** For Each Group, Immunoreactivity of NOS in CA1, CA2 and CA3 Regions of Hippocampus was Scored According to Five-Stage Grading System (x)

<table>
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<th></th>
<th>CA1</th>
<th>CA2</th>
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<tbody>
<tr>
<td>Group 1</td>
<td>0 1 2 3 4</td>
<td>0 1 2 3 4</td>
<td>0 1 2 3 4</td>
</tr>
<tr>
<td>Group 2</td>
<td>- 3 5 1 -</td>
<td>- 4 5 -</td>
<td>- 2 5 2 -</td>
</tr>
<tr>
<td>Group 3</td>
<td>4 5 1 -</td>
<td>1 6 1 2</td>
<td>- 6 4 -</td>
</tr>
<tr>
<td>Group 1 – Group 2</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
</tr>
<tr>
<td>Group 2 – Group 3</td>
<td>p &lt;0.05 *</td>
<td>p &gt;0.05</td>
<td>p &gt;0.05</td>
</tr>
<tr>
<td>Group 1 – Group 3</td>
<td>p &gt;0.05</td>
<td>p &lt;0.05 *</td>
<td>p &lt;0.05 *</td>
</tr>
</tbody>
</table>

0: No staining, 1: stain concentration under 25%, 2: stain concentration between 25%-50%, 3: stain concentration between 50%-75% 4: stain concentration over 75%. Statistical analyze of the data was done using Pearson’s chi-square test.

*All p-values less than 0.05 were considered to be statistically significant.
Figure 3: Hematoxilsen-Eosine stained section showing CA1, CA2 and CA3 hippocampal regions under light microscope (X4 magnification, scale shows 100 microns).

Figure 4: An example of Hematoxilsen-Eosine stained section showing CA1 area of a hydrocephalus-induced rat (X400 magnification, scale shows 100 microns). a: Protected cell body and nucleus. b: Degenerating neuron.

of excitotoxic injury through elevation of extracellular glutamate due to decrease in cerebral blood flow. According to current knowledge, excitotoxicity has been shown to be an important component of pathogenesis of traumatic and ischemic injury of central nervous system. In addition, further evidence suggesting the role of excitotoxicity in many of the chronic neurological disorders, some where “anti-excitotoxic treatment” was considered to become a therapeutic choice, is increasing (11).

Memantine has been well demonstrated to have protective effects on secondary neuronal injury due to ischemia or trauma of the central nervous system (4, 5, 9, 21, 22, 24). Regarding its neuroprotective effects on hippocampus, memantine has been shown to significantly decrease hippocampal neuronal cellular death emerged one week after cortical impact injury model in the rat (22). Memantine, at 20 mg/kg of daily dose, was reported to exhibit a significant neuroprotective effect in experimental models where transient or permanent focal cerebral ischemia has been induced (2, 22).

The purpose of this study was to evaluate whether anti-excitotoxic treatment could protect or attenuate the adverse effects of hydrocephalic process on hippocampal neurons. To our best knowledge, this study is the first to evaluate effects of anti-excitotoxic treatment on the hippocampal neurons in a rat model of infantile hydrocephalus. Considering the proposed significant role of extracellular glutamate elevation in the excitotoxic injury of hippocampal cells in hydrocephalus process, we preferred memantine (a non-competitive NMDA receptor antagonist) for anti-excitotoxic treatment in this study. Regarding the use of memantine in infantile hydrocephalus, there is only one study in the literature which, reported the effects of memantine treatment on esophageal and gastric smooth muscle reactivity (1). In that study, memantine treatment has been shown to influence the impaired contractile and relaxant activity of lower esophageal smooth muscle in hydrocephalus induced infantile rats (1). The present study is the first to report the effects of memantine treatment on hippocampal neurons in experimentally induced infantile hydrocephalus.

Our data showed that the average number of preserved neuronal cells with a clearly defined cell body and nucleus was significantly higher in CA1 and CA2 regions of the memantine treated hydrocephalic rat pups compared to non-treated hydrocephalic animals. In other words, daily administration of 20mg/kg memantine when started in the early stages of the hydrocephalic process, significantly prevented neuronal degeneration in CA1 and CA2 subunits of the infantile rat hippocampus in radiologically verified hydrocephalus. These findings seem to be supported by the immunohistochemistry data of the study revealing that the increase in NOS immunoreactivity was significantly reduced in CA1 and CA2 regions of the rats that had received memantine treatment during the hydrocephalic process compared with the non-treated rats. The data of the present study, which revealed the significant increase of NOS immunoreactivity in CA1, CA2 and CA3 regions of hippocampus in hydrocephalus induced rats, seem to be in agreement with the findings of Klinge et al. (16) who found a significant increase in hippocampal nitric oxide synthase activity two weeks after the induction of kaolin hydrocephalus. The increase in hippocampal NOS immunoreactivity, especially in the pyramidal layer of its CA1 subfield, is known to reflect the ischemia induced injury of the
hippocampal neurons (15). In experimentally induced adult hydrocephalus, the increase in hippocampal NOS activity has been suggested to be induced by increased extracellular glutamate levels and to imply a toxic neuronal response (16). In our model of kaolin-induced infantile hydrocephalus, the data about the effectiveness of memantine treatment in reducing NOS immunoreactivity in CA1 and CA2 regions of the hippocampus, seem to support the hypothesis that hydrocephalic process may affect hippocampal cells by the mechanism of excitotoxic injury through elevation of extracellular glutamate. The present findings imply the possibility of that the protective effect of memantine on hippocampal neurons might be mediated by its glutamate-NMDA receptor antagonism. A shortfall of the present study is the lack of cerebral blood flow measurements. Since simultaneous measurements of cerebral blood flow could not be performed, it was not possible to conclude whether or not this effect of memantine was mediated through any probable effect on cerebral blood flow. We are also far to interpret the data about the CA3 subunit of the hippocampus which seemed to have affected by the hydrocephalic process but have not displayed any significant changes both in neuronal count and NOS immunoreactivity parameters in memantine treated subjects. Further comprehensive studies are needed to clarify those points in a detailed way.

In conclusion, the findings of the present study simply point out that, hippocampal neurons may constitute important targets for injury secondary to hydrocephalic process, also in the infantile hydrocephalus model Early anti-excitotoxic treatment with memantine, a non-competitive NMDA receptor antagonist, seem to reduce or attenuate the adverse effects of hydrocephalic process especially in the CA1 and CA2 subunits of the hippocampus. These results appear to support the arguments justifying the presence of the mechanism of “glutamate associated excitotoxic insult” for hippocampal injury observed in experimentally induced hydrocephalus. We believe that the applicability of anti-excitotoxic treatment in preventing neuronal injury secondary to hydrocephalic process should be set forth with future comprehensive experimental and clinical studies.
REFERENCES


23. Socci DJ, Bjugstad KB, Jones HC, Pattisapu JV, Arendash GW: Evidence that oxidative stress is associated with the pathophysiology of inherited hydrocephalus in the H-Tx rat model. Exp Neurol 160:117–128, 1999
