Neuroprotective effects of tetracyclines on blunt head trauma: An experimental study on rats

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ABSTRACT

Background: Prevention of primary damage caused by head trauma may be avoided with protective measures and techniques which is a public health concern. Experimental and clinical studies about treatment of head trauma were all centered to prevent secondary damage caused by physiopathological changes following primary injury. Neuroprotective features of tetracyclines were the focus of several experimental studies in the last decade. In the present study we aimed to investigate the neuroprotective effects of tetracycline in an experimental model of blunt brain injury in rats. Materials and Methods: 32 male Sprague-Dawley rats were divided into four experimental groups (n = 8). Head trauma was not performed in control group (group 1, craniectomy only). In the second group, head trauma and craniectomy were performed. Intraperitoneal saline was used in addition to trauma and craniectomy for treatment in group 3 whereas intraperitoneal tetracycline and saline were used for treatment in group 4. Results: When histological examinations performed by transmission electron microscopy were evaluated, injury at ultrastructural level was demonstrated to be less pronounced in tetracycline group with decreased lipid peroxidation levels. Conclusion: In accordance with these findings, we conclude that systemic tetracycline administration is effective in reduction of secondary brain damage and brain edema and thus it may be considered as a therapeutic option.

Key words: Lipid peroxidation, neuroprotection, rat head trauma, secondary brain injury, tetracycline

Introduction

Traumatic brain injury usually involves an external impact targeting cranium and it is one of the leading causes of mortality and morbidity among young adults.¹ Head traumas in Turkey are usually associated with traffic accidents with an estimated mortality of 10/100,000. A thorough and detailed research focusing on head traumas is essential.²⁻⁵ The results of previous studies on neural trauma demonstrated two stages of injury defined as: Primary injury mainly linked to physical effect and secondary injury mainly associated with responses to primary effect.⁶⁻⁸

Tetracycline is an antimicrobial agent which has been shown to have neuroprotective effects including inhibition of caspase-1 and 3, nitric oxide synthase,
matrix metalloproteinases as well as glutamate excitotoxicity. So far, tetracyclines were proven to be beneficial against neuronal injury models. The enzymatic steps described in the above paragraph are important triggers for secondary injurious process which is involved in augmentation of primary injury. In the present study, we aim to investigate the neuroprotective effects of tetracycline on rat head trauma model of blunt cerebral injury in terms of mitochondrial and axonal degeneration as well as level of lipid peroxidation.

**Materials and Methods**

**Experimental design**

The study protocol was approved by the Ethic committee of Ankara Education and Research hospital. All experiments were performed at the research laboratory of the same center and specimens were evaluated at the Department of Histology of Hacettepe University Medical School.

32 male Sprague-Dawley rats were divided into four experimental groups as control, sham, SF and tetracycline group. Each group consists of eight rats and details of experimental protocol from each group are summarized in Table 1.

**Anesthesia and experimental procedure**

All rats were kept at standard conditions before and after experimental procedure. A mixture of xylazin (Rompun®, 2% solution, 60 mg/kg) and ketamine (Ketalar®, 5% solution, 10 mg/kg) was administered intraperitoneally for anesthesia. After immobilization of rats over stereotactic frame, a blunt head injury model was performed on rats of groups 2, 3 and 4 in accordance with the previous studies. Scalp was shaved and after local preparation with povidone iodine, a longitudinal incision of 2 cm was marked at the midline. Craniectomy was performed on left parietal region with a diameter 4 mm and dura was exposed for model of trauma [Figure 1a and b].

In the present study we investigated the neuroprotective efficiency of tetracycline on brain edema secondary to brain injury following blunt head trauma. A 24 gr weighted steel rod was dropped through a cylinder (attached to a plastic plate) from 10 cm above to the craniectomy region for blunt head trauma. Experimental site was closed and tetracycline hydrochloride was administered intraperitoneally at a dose of 90 mg/kg. Tetracycline was used at the same intraperitoneal dose as it was used in other studies (90 mg/kg). The rats were kept in cages at a room temperature of 24°C under standard animal care. At 24 hours, decapitation was completed after perfusion procedures and anesthetic overdose was given to rats for sacrifice with resection of whole cerebrum as single specimen [Figure 2]. Removed brain tissue was evaluated with quantitative lipid peroxidation measurements. Qualitative histological parameters which were converted to numerical data using USS system were developed from ultrastructural evaluation.

**Measurement of lipid peroxidation**

After homogenization of brain tissues a solution was prepared. The solution containing 10% brain tissue within 100NMKPO4 buffer (pH 7.3). Assessment of lipid peroxidation level was performed according to Uchiyama's thiobarbituric acid method. At least two graphical results obtained for each sample to determine lipid peroxidation levels as gram tissue/nanomol.

**Ultrastructural examination with transmission electron microscopy**

Brain tissue samples were fixed with 2.5% glutaraldehyde for 24 hours. The tissues were rinsed with Sorenson’s phosphate buffer, postfixation was performed with 1% osmium tetroxide. Following serial steps epoxy resin embedded sections were prepared TEM examination. The scores obtained from TEM examination were shown in accordance with ultrastructural parameters [Table 2].

**Statistical analysis**

Mann Whitney-U test was performed for group comparisons and P < 0.05 was accepted as significant.

| Table 1: Division of groups and procedure performed at each experimental group |
|---------------------------------|-----------------|
| Experimental Groups | Procedure |
| Group 1 | Craniectomy |
| Group 2 | Craniectomy+head trauma |
| Group 3 | Craniectomy+head trauma+physiological saline |
| Group 4 | Craniectomy+head trauma+physiological saline+tetracycline hydrochloride |

**Figure 1:** (a and b) Surgical incision and induction of head trauma
Results

Lipid peroxidation levels
Results of lipid peroxidation levels (nmol/g tissue) are shown in Table 3. The values are demonstrated as bar graphics in Graphic 1.

There was no significant difference between group 2 and 3 in terms of lipid peroxidation levels ($P > 0.05$). However, a comparison of the values of the tetracycline group (group 4) between the groups (Groups 2 and 3) designated above showed a statistical significance which proved tetracycline to be beneficial ($P < 0.05$).

Histological results
Transmission electron microscopic examination of samples and results of ultrastructural scoring system with averages for each group are demonstrated in Table 4. From each group four different samples were examined by blind investigators to reduce counting errors in each group. TEM images for each group are demonstrated in Figure 3a-d.

Efficacy of trauma was compared between C1 and C2 groups. A significant difference has been detected in terms of mitochondrial and axonal injury levels for these groups ($P > 0.05$). The systemic saline administrated group C3 and the control group C2 exhibited severe injury patterns such as myelin detachments and mitochondrial hydrops. The experimental group which received tetracycline exhibited minimal neuronal injury patterns. While intracellular vacuoles and mitochondrial hydrops were detectable in this group they were not

Table 2: Table includes scores of electron microscopic evaluation at ultrastructural level in terms of mitochondria, axonal morphology for normal and large myelinated axons. The evaluation was repeated for all experimental groups (100 counts were made for M (mitochondria), large and normal myelinated axons in each group

| Electron microscopic ultrastructural evaluation parameters | Mitochondria | Large myelinated axons | Normal myelinated axons |
|----------------------------------------------------------|--------------|------------------------|------------------------|
| Scoring                                                  | 0            | 1                      | 0                      |
|                                                         | Normal       | Cristas exposed        | Normal                 |
|                                                         | 1            | Hydroposed             | Detachment of myelin strata |
|                                                         | 2            | Amorph mass exposed    | Interruption of myelin strata |
|                                                         | 3            |                        | Honeycomb appearance    |

Figure 2: Resected whole brain specimen after perfusion procedures and decapitation

Figure 3: (a) Transitional electron microscopic (TEM) image of group 1 showed minimal detachment of myelin strata, arrow shows mitochondria. (b) Detachment of myelin strata (multiple arrows). (c) Similar ultrastructural alterations with prominent detachment of myelin strata (arrows). (d) Minimal ultrastructural alteration compared to findings of groups 2 and 3

Graphic 1: Lipid peroxidation measurements (MDA levels) in all experimental groups (MDA levels K1: Control, K2: Trauma, K3: Saline, D: Tetracycline)
Despite tremendous number of experimental and clinical trials so far, a considerable and cost-effective pharmacological agent to prevent secondary brain injury was not able to be found. A substantial increase in intracranial pressure and development of hypoxia at the early stages of trauma is directly proportional with a rise in mortality and morbidity.\textsuperscript{[23,24]}

Increase in excitatory amino acids (EAA) exists as an important mechanism after head injury and is the major concern of several studies.\textsuperscript{[27,28]} Free radical formation is triggered by excitatory amino acids and lipid peroxidation are accepted as the most important mechanisms that cause secondary brain damage.\textsuperscript{[8,29]}

Table 3: Lipid peroxidation values in nmol/gram tissue for each experimental group

| Experimental Groups | Lipid peroxidation (nmol/gram tissue) |
|---------------------|---------------------------------------|
| Group 1             | 48,681±11,123                         |
| Group 2             | 66,681±8,585                          |
| Group 3             | 65,712±3,049                          |
| Group 4             | 55,106±7,992                          |

Table 4: Numerical values regarding 100 counts obtained from each experimental group in terms of mitochondria, normal myelinated axons and large myelinated axons. Average values were shown under the last row of each section. Four different samples were examined for each group by blind investigators to reduce counting errors in each group

| Mitochondria types | Group 1 | Group 2 | Group 3 | Group 4 |
|--------------------|---------|---------|---------|---------|
| Mitochondria sample 1 | 11      | 33      | 27      | 25      |
| Mitochondria sample 2 | 10      | 31      | 26      | 23      |
| Mitochondria sample 3 | 9       | 32      | 28      | 26      |
| Mitochondria sample 4 | 12      | 34      | 26      | 22      |
| Average values      | 10,50   | 32,50   | 26,75   | 24      |
| Large myelinated axons sample 1 | 8       | 35      | 35      | 32      |
| Large myelinated axons sample 2 | 6       | 39      | 37      | 29      |
| Large myelinated axons sample 3 | 7       | 36      | 39      | 25      |
| Large myelinated axons sample 4 | 5       | 39      | 37      | 22      |
| Average values      | 6,50    | 37,25   | 37      | 27      |
| Normal myelinated axons sample 1 | 0       | 26      | 19      | 16      |
| Normal myelinated axons sample 2 | 1       | 20      | 24      | 14      |
| Normal myelinated axons sample 3 | 0       | 24      | 17      | 12      |
| Normal myelinated axons sample 4 | 0       | 28      | 22      | 14      |
| Average values      | 0,25    | 24,50   | 20,50   | 14,00   |

as severe as encountered in C2 and C3 groups. The difference was significant favoring a neuroprotection for experimental samples ($P < 0.05$).

Discussion

Tetracycline is a well-known antibiotic proved to have protective role in neural injury. In the present study, we investigate its neuroprotective role in experimental head trauma. We found that tetracycline reduced the lipid peroxidation levels and findings on electron microscopy in experimental head trauma.

52,000 yearly deaths and 80,000 permanent severe cerebral injury are reported in the USA per year.\textsuperscript{[2,3]} Despite the limited number of statistical studies about the frequency of head traumas in Turkey, traffic accidents are reported to be the leading cause of head trauma in Turkey. 60% of deaths due to head trauma originated from traffic accidents and a health problem which has a high mortality and morbidity.\textsuperscript{[23,24]}

When the head is subjected to a severe trauma, a direct damage (primary tissue damage) to cortical tissue is noted. Several mechanisms are involved in primary brain injury such as direct trauma to brain parenchyma, smashing of brain tissue by bony protuberances and penetration of brain tissue by bone fragments or foreign bodies. Skull fractures, intracranial bleeding, coup and countercoup lesion, concussion and diffuse axonal injury are the consequences of primary lesion and secondary tissue damage usually follows the primary.\textsuperscript{[25]} Prevention of primary damage caused by head trauma may only be accomplished by preventive measures and is a subject of public health concern.\textsuperscript{[26]}

Intracranial pressure rises after head injury which might interfere with critical level of cerebral blood flow. This reduction in blood flow further exacerbates the secondary injury process.\textsuperscript{[8]} Secondary brain injury involves several mechanisms mainly associated with metabolic responses and effects of mechanical pressure.\textsuperscript{[28]} Current scope of experimental and clinical studies about treatment of head trauma were to prevent secondary damage caused by pathophysiological cascades following primary injury.\textsuperscript{[7,8]} In the present study we aimed to prevent the secondary injury in experimental head trauma.

Tetracycline is a well-known antibiotic and its neuroprotective role was proven in several animal models within the last decade.\textsuperscript{[13,21,33‑36]} Tetracyclines can easily cross blood–brain barrier due its hydrophilic properties. Its mechanism is associated with several pathways that
results in reduction of apoptosis, neuroinflammation, infarct size, and vascular injury.[35,37] Koistinaho, et al. reported tetracyclines to be a neuroprotective agent for brain ischemia in the rats as well as cerebral ischemia, Parkinson’s disease, Huntington’s disease, amyotrophic lateral sclerosis and multiple sclerosis.[33,36] The study performed by Kraus, et al. points to protective role of minocycline on neuronal cultures.[38] In the present study we investigate the use of tetracycline on rat head trauma. A comparison of the trauma group and the tetracycline-treated group demonstrated a neuroprotective effect of drug on head trauma.

In the present study a comparison of lipid peroxidation levels between head trauma and tetracycline group did not reveal any significance designating an overt neuroprotective role of tetracycline in head trauma. Furthermore, electron microscopic examinations showed less prominent findings in the experimental head trauma group.

### Conclusion

Tetracycline was shown to be beneficial in blunt head trauma when lipid peroxidation and histology of the present study was considered. We conclude that systemic tetracycline may be a considerable therapeutic option since its administration may ameliorate findings of secondary brain damage and edema.

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