ORIGINAL RESEARCH

Correlation Between Rats Hippocampal ATP Content and Delayed Encephalopathy After Acute Carbon Monoxide Poisoning

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Objective: To study the correlation between the adenosine triphosphate (ATP) content in the hippocampus of rats and delayed encephalopathy after acute carbon monoxide (CO) poisoning.

Methods: A total of 40 male Wistar rats weighing 180–230g, in accordance with the random number table, were selected and divided into the delayed encephalopathy after acute carbon monoxide poisoning (DEACMP: Rats with cognitive impairment after carbon monoxide poisoning) group (n = 32) and the control group (n = 8). A DEACMP rat model was generated by inhalation of CO. The Morris water maze evaluated the ability to learn and memorize in rats. The changes in neurons in the hippocampus of the rats were observed by hematoxylin-eosin (HE) staining. Lastly, the ATP content in the hippocampus of the rats was measured by high-performance liquid chromatography (HPLC).

Results: The ATP content of the experimental group was significantly higher than that of the control group in the hippocampus of the rat model, so the difference was statistically significant (P < 0.05); the intra-group comparison was made for the ATP content in the experimental group, and the difference was statistically significant as group 21d > group 14d > group 7d (P < 0.05); and no significant difference was found between group 21d and group 28d (P > 0.05).

Conclusion: The changes in the ATP content in the hippocampus of the rats are correlated with the occurrence of delayed encephalopathy after acute carbon monoxide poisoning; it may take part in the pathogenesis of DEACMP. This offers some elicitation to the prevention and treatment of the disease.

Keywords: carbon monoxide poisoning, delayed encephalopathy, ATP, HPLC

Introduction

Usually, delayed encephalopathy after acute carbon monoxide poisoning (DEACMP) is considered a type of central nerve injury. DEACMP is the most common and serious complication of acute carbon monoxide (CO) poisoning,¹ which results in a high disability rate and fatality rate and thus seriously affects the quality of life of patients. To date, the pathogenesis of DEACMP remains unclear, and it is difficult to cure due to the lack of an effective treatment method and strategy. This is a difficult problem in urgent need of a clinical solution.²

Adenosine triphosphate (ATP) is not only a classical energy substance, but also a neuromodulator and neurotransmitter, and its receptors are widely distributed in the nervous system. Studies³ have confirmed that after partial injury of central nervous system, high concentration of ATP accumulates in the surrounding area and participates in secondary apoptosis or necrosis of nerve cells. Other scholars^{4,5} have also reported the effect of ATP on spinal cord neurons of cultured embryonic mice. At present, the role of ATP and its receptors in the occurrence of nervous system diseases has become a new research hotspot. Therefore, we investigated whether ATP is related to the DEACMP.

In order to elucidate the pathogenesis of DEACMP and explore new therapy strategies, an optimal method was selected to generate the model of delayed encephalopathy after acute CO poisoning by exposure to 3000 ppm CO for 40 min, based on the reports of Hara et al⁶ that featured low mortality and high incidence of DEACMP. For the DEACMP rat model, the Morris water maze was employed to collect the average escape latency, frequency of passing the platform and other parameters to evaluate the ability of learning and memory in rats. In addition, high-performance liquid chromatography (HPLC) was applied to measure the change of ATP content in the hippocampus of the DEACMP rats. An exploration of the correlation between ATP content in hippocampus of rats and delayed encephalopathy provides a foundation for further research of the pathogenesis of delayed encephalopathy.

Experimental Material

Laboratory Animal

Forty healthy male Wistar rats, specific-pathogen-free grade (SPF), weighing 180–230g (Research Center for Laboratory Animal Science Inner Mongolia University, approval No. SCXK [M] 2016-0001), were fed in clean grade during the experiment. Before the experiment, they were given adaptive feeding with standard feed and ad libitum access to water for 1 week; they were maintained in a 12-hour light/dark cycle with cage temperature at 18–23°C.

Main Reagents and Equipment

The instruments and reagents used for this study included a CO exposure box (manufactured by the Hyperbaric Oxygenation Department of Baotou Central Hospital, Inner Mongolia), high-performance liquid chromatograph (Thermo UltiMate 3000), ODS C18 chromatographic column (Thermo), refrigerated centrifuge (Thermo), ultraviolet spectrophotometer (T6 Xinyue), LEICA CM1950 cryostat, LEICA DM 2500 microscope, homogenizer (JY98-III), pure CO gas (99.99%, manufactured by Foshan KODI Gas Chemical Co., Ltd.), ATP standard (Sigma-Aldrich) stock solution prepared with ultrapure water, and other reagents in the experiment that were all domestic chromatographic or analytical reagents.

Experimental Method

Morris Water Maze Training

After one week of adaptive feeding, the laboratory animals were labeled for water maze training. The laboratory rats were put into the water with their heads facing the tank wall at the middle points of the four quadrants, respectively. The path and time of the rats to find the platform in the water were observed and recorded. If any rats failed to find the platform within 120s, they were guided to it. The rats were left on the platform for 30s. The rats were trained 2–3 times per day. After 5 days, the path represented the tendency in a straight line to the platform. The rats with escape latency less than 1 min were recorded as qualified, and those with escape latency greater than 1 min were eliminated.

Animal Grouping

The qualified rats were randomly divided into 2 groups, namely the DEACMP group and the control group.

Establishment of DEACMP Animal Model

The rats in the experimental group were randomly placed in a CO exposure box with a concentration of 3000 ppm for 40 min to observe the toxic reaction of the animals. Those in the control group were placed in an air-filled exposure box.

Blood HbCO Titer

The carboxyhemoglobin (HbCO) titer in rats was measured at 15 min, 30 min, 1 h and 2 h after poisoning. The modified dual-wavelength quantitation method⁷ was used to measure the concentration of HbCO in the rat tail-vein blood in each group. Blood was collected from the tail vein. First, 0.1 mL of rat tail-vein blood was added to 20 mL of 0.4 mol/L

HbCO (%)= $(2.44 \times A_{535}/A_{578}-2.68) \times 100\%$

Measurement of ATP Content by High-Performance Liquid Chromatograph

According to the experimental methods,^{8–11} the ODS C18 chromatographic column (4.6 mm \times 250 mm, 5 um) was utilized. Mobile phase: 99% 0.18 mmol/L phosphate buffer solution (28g KH₂PO₄ was dissolved in 1145 mL of distilled water, and filtered with 0.45 mml filter membrane before use, PH = 6.25), 1% pure methanol; detection wavelength: UV wavelength 254 nm; sample size: 10 uL; column temperature: room temperature.

Thirty-two surviving rats after CO poisoning (n = 32) were randomly divided into 4 groups (7d, 14d, 21d, and 28d, n=8/group). Chloral hydrate (10%) was administered to each group via intraperitoneal injection (0.3 mL/100 g) for anesthesia, after which the rats were decapitated, and the skulls were opened to allow the removal of the brain. The brains were placed on the ice plate, and the meningeal blood vessels were removed to an ice table for the hippocampus tissues. The tissues were weighed, respectively, and added with 0.6 M cold perchloric acid solution (85.8 mL perchloric acid was dissolved in 1638.1 mL distilled water and then filtered with 0.22 um filter membrane) to make a 10% homogenate (by mechanical and ultrasonic homogenizing), which was centrifuged at 1000 r/min for 20 min at low temperature. Then, 0.4 mL of the supernatant was added to the 3 mol/L K₂CO₃ solution to adjust the pH to neutral, centrifuging at 3000 r/min at 4°C for 5 min. Afterward, the supernatant passed through a 0.22 um filter membrane to obtain a sample size of 10 uL.

Preparation of Standard Curve

The ATP standard was accurately weighed to prepare 0, 40, 80, 120, 160, and 200 μ g/mL of ATP standard solution, which was added with 10 uL of sample into the chromatograph, respectively, under the above chromatographic conditions. A regression of mass concentration vs peak area was performed to each group to obtain the standard curve equation.

Statistical Analysis

All data analyses were performed using the statistical software SPSS 20.0. Quantitative data are expressed as the mean \pm standard deviation; one-way ANOVA was used to compare quantitative data between groups; the *t*-test was employed for pairwise comparisons between groups; and a **P* < 0.05 was considered statistically significant.

Results

Identification of Animal Models

After inhalation of CO gas, the rats in the experimental group presented indications of hyperactivity and dysphoria after about 15–20 min, and their toes and mucous membranes appeared pink in color. After 20 min, the rats showed signs of shortness of breath, limb paralysis, and convulsions; opisthotonus occurred successively, and some rats died. No such manifestation was observed in the control group. The HbCO levels in the rats' tail-vein blood after poisoning were measured at multiple time points (Table 1). The results showed that the rats were poisoned according to the COP standard.¹² The difference between the two groups was statistically significant (*P < 0.05).

Water Maze Test Results

The data were collected by the Morris water maze, and the results showed that the memory of the rats after poisoning was impaired. Specifically, the average latency increased, and the frequency of passing the platform decreased. The frequency of passing the platform of fewer than 3 times is indicative of DEACMP (Figure 1B).^{13,14} However, no obvious change was observed in the control group (Figure 1A, <u>Additional Table</u>).

Group I 5min 30min 60min I 20min Experimental group 46.17±12.24 20.90±2.68 9.58±4.66 3.53±2.12 Control group 2.40±1.56 2.56±1.32 2.30±1.17 2.46±0.94 *P (compared with control group) *P < 0.05 *P < 0.05 *P < 0.05 *P < 0.05					
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	Experimental group Control group *P (compared with control group)	46.17±12.24 2.40±1.56 *P < 0.05	20.90±2.68 2.56±1.32 *P < 0.05	9.58±4.66 2.30±1.17 *P < 0.05	3.53±2.12 2.46±0.94 *P < 0.05

 Table I HbCO Concentration of Rat Tail-Vein Blood at Each Time Point (%)

Note: *P is compared with the control group.

Pathological Changes of Hippocampus of Rats

The pathological sections of rat brain tissue were stained with hematoxylin and eosin. Under the ordinary optical microscope, it can be seen that the hippocampal cells in the control group were plump, the perinuclear pale circle was obvious, and the cells were arranged orderly (Figure 2A). In the rats with delayed encephalopathy (Figure 2B), extensive damage to cells decreased cell number, thinning of the cell layer, disorder of cell structure, obvious cell shrinkage, pyknosis, and hyperchromatic nuclei, and unclear structure was observed in the hippocampus and subcortex under a microscope. However, no obvious focal necrosis was observed.

Establishment of ATP Standard Curve

The peak time of the ATP standard was measured to be 5.278 min (Figure 3). The peak values of 0, 40, 80, 120, 160, and 200 ug/mL ATP standard solution were measured, respectively, and the measured stable peak time was 5.278 min. The linear relationship was obtained via the proportional relationship, and the standard curve Y (peak area) = 0.2492X (ATP content) – 0.6363 (Figure 4).

Results of Sample Determination

The prepared samples were analyzed, and the ATP chromatogram of the rat hippocampus was obtained (Figure 5). Compared with the control group, the ATP content of the delayed encephalopathy group model in the experimental group significantly increased, and the difference was statistically significant (*P < 0.05). The intra-group comparison was performed for the ATP content of the DEACMP group. The results showed group 21d > group 14d > group 7d, and the difference was statistically significant (*P < 0.05 is compared with 7d, & P < 0.05 is compared with 14d). The difference between group 21d and group 28d was not significant (*P > 0.05, Table 2). The results showed that hippocampal ATP in



Figure I Morris water maze experiment (spatial exploration experiment), the red circle in the figure indicates the position of platform. (A) Rats of the control group. (B) DEACMP rats.



Figure 2 (A) Hippocampus of rats in the control group (HE). (B) Hippocampus of rats in the experimental group (HE).



Figure 3 Chromatogram of ATP standard.

delayed encephalopathy rats was higher than that in normal rats, proving that hippocampal ATP content was related to DEACMP, and we have added the relative description in the Results.

Discussion

The pathogenesis of DEACMP is not presently clear. The establishment of an accurate and reliable animal model is the basis for further experimental research. At present, there are two main modeling methods: intraperitoneal injection and inhalation. The former features simple operation, strong controllability, and experimental gas savings. However, this method is different from the normal poisoning method in that the CO absorption method is different, and the poisoning time arising out of this method is not continuous. As a result, the amount of CO absorption is not constant and can cause many deaths in laboratory animals, which can then affect the experimental results. The CO inhalation method is more



Figure 4 ATP standard curve.



Figure 5 Chromatogram of sample for hippocampus of rats.

similar to that of clinical patient poisoning in that the toxic gas enters the blood in the same way. In this sense, an optimal modeling method was selected to generate the DEACMP model by the inhalation of 3000 ppm CO gas for 40 min, according to the reports of Hara et al. After the model was established, the Morris water maze test was performed for the

Group	ATP (ug/mg×10-2)		
Normal control group	4.176±1.169		
Poisoning group 7d	9.824±2.839 (*P < 0.05)		
Poisoning group 14d	11.642±3.443 (*P < 0.05, [#] P< 0.05)		
Poisoning group 21d	13.872±4.317 (*P < 0.05, ^{&} P< 0.05)		
Poisoning group 28d	13.561±4.120 (^{\$} P > 0.05)		

Table 2 ATP Content in Hippocampus of Rats in DifferentExperimental Groups

Note: **P* is compared with the control group, ${}^{\#}P$ is compared with 7d, ${}^{\&}P$ is compared with 14d, and ${}^{\$}P$ is compared with 21d.

experimental group, and the results showed that the rats with delayed encephalopathy exhibited abnormal behavior, such as increased latency and decreased frequency of passing the platform; HE staining was used to observe the pathological changes in the hippocampus of the rats. In the experimental group, the pathological manifestations of delayed encephalopathy appeared in the hippocampal area. The experimental results showed DEACMP manifestations, proving that the animal model established in this experimental study is ideal and reliable.

The pathogenesis of DEACMP is currently believed to be correlated with the apoptosis mechanism, the theory of ischemia and hypoxia, the inflammatory injury mechanism, the immune injury mechanism, and the CO/NO signal transduction mechanism.^{15–18} Some studies³ have confirmed that high concentrations of ATP accumulate in the surrounding area after partial central nerve injury and participate in the apoptosis or necrosis of secondary nerve cells. ATP is not only a classical energy substance but also a neuromodulator and neurotransmitter. Its receptors are widely distributed in the nervous system. At present, the ATP and its receptors have become a new research hotspot, and the role of ATP in the occurrence of neurological diseases has also drawn the attention of scholars. Studies by other scholars^{4,5} reported the effect of ATP on spinal cord neurons of cultured embryonic mice. The results showed that low concentrations of ATP could increase the activity of cells, while high concentrations of ATP could reduce the activity of cells and inhibit the formation of neuronal processes. For the experimental group, the precise scientific HPLC method was employed to measure the ATP content in the hippocampus of rats in different time periods in order to explore whether the ATP content in the hippocampus is correlated with the occurrence of delayed encephalopathy.

The results of this study showed that the ATP content in the hippocampus of DEACMP rats increased significantly, especially in the period from 14 to 21 days. This suggests that the ATP content in the hippocampus of rats is correlated with the occurrence of delayed encephalopathy. Through this experiment, it was clear that the content of ATP in the nervous system of rats with delayed onset encephalopathy was significantly increased, and it was also proved that delayed onset encephalopathy would affect the energy metabolism of nerve cells. However, it remains to be further studied whether ATP is increased through what pathway and whether ATP plays a protective or destructive role in the pathogenesis of tardive encephalopathy. This study was only based on the analysis of the changes in ATP content in nerve cells of rats with toxic tardive encephalopathy. The lack of specific reasons for the changes of neurotransmitters in neurons and the change principle of energy pathways requires further research and analysis. The results of this study can provide research basis for the pathogenesis and further diagnosis and treatment of delayed onset encephalopathy.

Ethical Approval

All experiments were evaluated and approved by the ethics Committee of Hulunbuir people's Hospital and complied with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

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Role and mechanism of P2Y12 receptor-mediated microglia activation in delayed encephalopathy caused by acute carbon monoxide poisoning(81641040).

Disclosure

The authors declare that they have no competing interests in this work.

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