Effect of methylprednisolone on the axonal impairment accompanying cellular brain oedema induced by water intoxication in rats

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Abstract

OBJECTIVES: Our previous experiments proved that methylprednisolone (MP) can significantly reduce axonal impairment accompanying extracellular oedema induced by the osmotic challenge (load) on the blood-brain barrier (BBB). The aim of the present work was to identify whether MP can affect myelin impairment accompanying intracellular oedema induced by water intoxication.

METHODS: For induction of cellular brain oedema, the standard model of water intoxication was chosen. Animals received distilled water in amount corresponding to 15% of the animal’s body weight. The volume was divided into three parts and administered intraperitoneally in 8 hours interval. Axonal changes were recognized as signs of myelin disintegration (oedematous distensions, axonal swelling, vesicles, varicosities) at histological sections stained with Black Gold and classified into four grades of myelin degradation. Hippocampal CA1 and CA3 areas and the dentate gyrus were selected for the study. Methylprednisolone was administered either intraperitoneally or intracarotically. Its effect was studied in two different time intervals: in the acute group (30 minutes after hyperhydration and MP application) and in chronic one (1 week after hyperhydration and MP application).

RESULTS: In both the acute and chronic groups, cellular oedema induced by water intoxication brought about apparent damage of myelin (compared to control animals p<0.0001). Intracarotical injection of MP was not able to influence myelin integrity changes either in the acute or in chronic group. However, intraperitoneal administration of MP increased the level of myelin deterioration in the acute group (p 0.05), but improved myelin changes in the chronic group (p<0.005).

CONCLUSION: The effect of MP on axonal impairment during cellular brain oedema induced by water intoxication differs from that during the extracellular osmotic oedema. In the extracellular oedema, cellular metabolism is not significantly affected and myelin changes can be influenced by the neuroprotective effect of MP. The primary cause of cellular oedema is a disorder of cellular metabolism and myelin impairment is one of the structural consequences of such disorder. That is why the myelin changes are not affected by MP administration in a consistent and specific manner.
INTRODUCTION

In our previous work we have shown that effect of methylprednisolone (MP) on axonal damage in an experimental model of cerebral oedema induced by extracellular osmotic insult to blood-brain barrier by mannitol was constant and specific. For a constant effect of MP we considered the fact that the MP in both time intervals and in both modes of administration the degree of myelin disintegration. The specific effect of MP means that in both time intervals the degree of myelin impairment was reduced (Kozler et al. 2011). In the present study the effect of MP on axonal damage was followed in an experimental model of cellular brain oedema induced by water intoxication.

During water intoxication (hyperhydration) an osmotic gradient, is formed which moves water into the intracellular compartment – a cellular oedema is induced (Go 1997; Kimberg 1995). Animal is hyperhydrated by intraperitoneal injection of distilled water in the total amount of 15% of the body weight. The total dose is divided into three portions, which are applied in eight-hour intervals over 24 hours. This method of induction of cellular oedema does not cause any deterioration of neurological status or an increased mortality, as is the case of acute intoxication, when the total dose is administered all at once. Method is a currently used as a standard experimental model of cellular brain oedema (Olson et al. 1994; Vajda et al. 2000; Manley et al. 2000; Yamaguchi et al. 1997).

Neuroprotective effect of MP was tested in experimental models of cerebral oedema with mixed results (Ildan et al. 1995; Park 1998; Shapiro et al. 1992; Lin et al. 1994; Slivka & Murphy 2001). MP has a special position in the management of CNS injuries. While in the treatment of spinal cord injury it is irreplaceable (Bracken et al. 1997), the brain injury does not apply to the absence of documented neuroprotective effect (Marshall 2000; Czekajlo & Milbrandt 2005; Edwards et al. 2005; Alderson & Roberts 1997).

Methylprednisolone is a synthetic steroid with four times higher glucotropic and one fifth of mineralocortic action of cortisol (hydrocortisone). The primary neuroprotective effect of MP, mechanisms by which MP affect both healthy and damaged CNS, and pharmacokinetics of MP including its penetration through the BBB has been described in detail in the literature (Hall 1992; Park 1998).

MATERIAL AND METHODS

For the experiment, adult animals of both sexes of the Wistar strain of laboratory rats (weight 350–450 g) were used. Animals were treated in accordance with the current Guidelines for the treatment of laboratory animals (EU Guidelines 86/609/EEC).

Induction of cellular oedema – water intoxication

Animals were hyperhydrated with the method of water intoxication (Go 1997; Manley et al. 2000; Marmarou et al. 2006; Yamaguchi et al. 1997). Each was given distilled water in a quantity equal to 15% of the body weight in three separate doses applied intraperitoneally at 8-hour intervals during 24 hours prior to launching the experiment.

Application of methylprednisolone

To study neuroprotective effect of methylprednisolone two experimental groups were formed: group A (acute) and group C (chronic). In each group three different experiments were performed, each using five animals. Results were compared with the control group (CG) of intact animals. In group A the following experiments were done: A1 – induction of oedema by water intoxication, A2 – induction of oedema by water intoxication accompanied with MP i.p. administration (the total dose of MP 100mg/kg was divided into three sub-doses; each sub-dose was administered together with the dose of distilled water during oedema induction). Perfusion and fixation (see below) were performed 30 minutes after the last water and MP administration. A3 – induction of oedema by water intoxication accompanied with intracarotid MP administration (in a single dose of 100mg/kg, 10 minutes after BBB opening by mannitol (see below), one hour after the last dose of distilled water). Perfusion and fixation were performed 30 minutes after the MP administration. Experiments in group C followed the arrangement of subgroups in group A, but the perfusion and fixation were performed 1 week after the last water and MP administration. The results were subjected to statistical analysis (see below).

Microsurgical exposure of the internal carotid (ACI)

Animals were put into the state of general anaesthesia using intraperitoneal application of thiopental in the dose of 4mg/100g and allowed to ventilate spontaneously throughout the procedure. Starting from a skin incision along the midline between the upper end of the sternum and the mandible, the whole common carotid artery (ACC, arteria carotis communis) was exposed with a standard microsurgical technique and, before its bifurcation, also the proximal portions of the internal carotid (ACI, arteria carotis interna) and external carotid (ACE, arteria carotis externa), which was ligated close beyond the bifurcation. An intraluminal catheter was introduced into the ACC trunk from the arteriotomy for selective application of mannitol. With
the application over and the catheter removed, the ACC was ligated distal to and proximal to the arteriotomy. The operation concluded with a single-layer suture (Kozler et al. 2011).

**Osmotic opening of the BBB + MP administration**

Mannitol 20% (200 g in 1 000 ml of water for injection, 1 098 mosmol/l) in a dose of 5ml/kg was selectively applied in the ACI at a rate of 0.12 ml/sec (Saris et al. 1988; Rapoport 2000).

In experiments A3 and C3, 10 minutes after Manitol injection, MP was administered intracarotically in the dose of 100mg/kg. After the surgical intervention, animals were placed in boxes offering standard access to food and drink.

**Perfusion and fixation**

Animals were sacrificed in a deep anaesthesia via standard transcardial perfusion with a 4% solution of paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) for 15 minutes. After removal from the skull, the brain was fixed in the same solution for 24 hours. Serial coronary sections (30 μm thick) were cut at vibratome from each brain, placed on gelatine-coated slides and dried.

**Neurohistology**

The sections were rehydrated and axonal changes detected with the Black Gold II method of staining (Histo-Chem Inc., Jefferson, AZ, USA.) (Schmued et al. 1999). The hippocampal formation was the main part of the brain under study because of its known high sensitivity to various pathogenic stimuli. Analysis was centred on the CA1 and CA3 areas of the hippocampus and on the dorsal blade of the dentate gyrus (DG).

The neurohistological picture of the structural integrity of the axons was assessed with the aid of the following grades of myelin degradation: 1 = no change, 2 = sporadic oedematous vesicles and sporadic oedematous axons, 3 = multiple vesicles, varicosity, oedematous axons and helical course of axons, 4 = myelin fragmentation.

Neurohistological examination was done in both experimental groups (consisting from 3 different experiments) and one control group, with five animals in each group. For the statistical analysis data from 10 sections of each brain were used.

**Statistical analysis**

The results were statistically evaluated using the t test and one-way analysis of variance (ANOVA) followed by Dunnett post hoc analysis. The statistical software GraphPad Prism version 4.00 for Windows (GraphPad Software, San Diego, CA, USA) was used for all statistical analyses.

**RESULTS**

It was shown in the previous paper, that brain oedema is accompanied with signs of myelin degradation. They include oedematous distensions, axonal swelling, vesicles, varicosities, helical course of axons, and myelin fragmentation (Kozler et al. 2011).

Results obtained in both experimental groups are shown at Figures 1 (acute group) and 2 (chronic group). In all experimental animals the cellular oedema induced by water intoxication brought about myelin impairment ($p<0.0001$). Such impairment was more serious in the chronic group (average grade 3.3) than in the acute group (average grade 2.7).

In the acute group, intraperitoneal MP administration worsened the degree of myelin disintegration ($p<0.05$); however intracarotid MP injection did not affect the process of myelin disintegration (Figure 1).
In the chronic group, intraperitoneal MP administration improved the signs of myelin disintegration ($p<0.005$), but intracarotid MP injection had no affect on the process of myelin disintegration (Figure 2).

**DISCUSSION**

Prevention of adverse effects of brain tissue oedema is the aim of many research groups. Promising appear to be substances with antioxidant effect (Gasparova et al. 2009). The widely used neuroprotective agent methylprednisolone (MP) reveals also significant antioxidative capacity (Park 1998).

In our previous study, we demonstrated the constant and specific effects of MP at the structural integrity of myelin impaired by extracellular brain oedema induced by osmotic challenges to blood-brain barrier (Kozler et al. 2011). The aim of the present study was to evaluate the effect of MP on myelin impairment resulting from cellular oedema induced by water intoxication. Results of the present study showed that cellular oedema can bring myelin disintegration and that these changes are time dependent – the most serious degree of disintegration was found one week after the oedema induction.

Similar experience had also other authors (Creed et al. 2011; Onaya 2002; Stys 1998). Our present results also showed that MP injected into the internal carotid artery had no effect on the myelin disintegration induced by cellular oedema both in acute and chronic groups. Intraperitoneal administration of MP however, further impaired the integrity of myelin in the acute group, but improved it in the chronic group ($p<0.005$) (Figures 1 and 2). The effect of MP can be thus regarded as non-constant, since its effect was inconsistent in both time-defined groups, and after both administration procedures. Effect of MP can be also called nonspecific, because it influences myelin disintegration in different ways – intracarotid administration brought no effect hence intraperitoneal administration resulted either in further myelin deterioration or in the integrity improvement.

In the present experiments, effect of MP on myelin impairment induced by cellular oedema was totally different from its constant and specific effects induced by extracellular oedema, which was published earlier (Kozler et al. 2011). Variability in the effect of MP can be related to different types and pathogenetic mechanisms of brain oedemas. Osmotic BBB insult leads to extracellular (vasogenic) oedema caused by the increased permeability of BBB, with the cellular metabolism primarily unharmed. Water intoxication induces cellular (cytotoxic) oedema where the primary cause results from damage to cellular metabolism with a relatively preserved selective permeability of BBB (Kimlberg 1995). In the primary cause of cellular oedema – in the impairment of the cellular metabolism – we see the cause of unreliability of MP effect on myelin disintegration. Onaya (Onaya 2002) on the basis of neuro-pathological studies had proved that axonal disintegration accompanying diffuse brain injuries is always preceded by cellular oedema and he concluded that the occurrence of damaged axons is not possible without cellular oedema. These findings in the context of results from our experimental studies allow concluding that the myelin damage resulting from cellular oedema probably cannot be influenced by MP, or the effect is not constant and specific as it was observed after the extracellular oedema. International clinical studies “CRASH trial” in which randomized patients with brain injury received intravenous infusion of MP had to be prematurely terminated because the outcome of patients who received MP was worse than in those who did not receive MP (Edwards et al. 2005; Czekajlo & Milbrandt 2005). Though only speculatively, we must conclude that our experimental results correspond with the clinical study, where the improvement of the clinical status of patients after MP administration was expected.

**CONCLUSION**

Results of our experiments show that effects of methylprednisone on the myelin impairment accompanying cellular brain oedema induced by water intoxication are neither consistent nor specific. This finding does not correspond with our previous findings that MP can significantly reduce the degree of myelin damage resulting from extracellular oedema induced by osmotic insult. The explanation for this contradiction lies in the different types of oedema, which lead to myelin damage. The extracellular oedema does not significantly affect the cell metabolism. Resulting changes in the myelin integrity are therefore more easily influenced by the documented neuroprotective effect of MP.

The primary cause of cellular oedema is a disorder in cellular metabolism which results, inter alia, in various structural changes, including myelin disintegration. Because between the cellular oedema and myelin damage a causal relationship was proven, it is very likely that methylprednisone does not affect myelin disorders related to cellular oedema either constantly or specifically.

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**REFERENCE**


