Evaluation of two experimental models of hepatic encephalopathy in rats

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Abstract

The serious neuropsychological repercussions of hepatic encephalopathy have led to the creation of several experimental models in order to better understand the pathogenesis of the disease. In the present investigation, two possible causes of hepatic encephalopathy, cholestasis and portal hypertension, were chosen to study the behavioral impairments caused by the disease using an object recognition task. This working memory test is based on a paradigm of spontaneous delayed non-matching to sample and was performed 60 days after surgery. Male Wistar rats (225-250 g) were divided into three groups: two experimental groups, microsurgical cholestasis (N = 20) and extrahepatic portal hypertension (N = 20), and a control group (N = 20). A mild alteration of the recognition memory occurred in rats with cholestasis compared to control rats and portal hypertensive rats. The latter group showed the poorest performance on the basis of the behavioral indexes tested. In particular, only the control group spent significantly more time exploring novel objects compared to familiar ones (P < 0.001). In addition, the portal hypertension group spent the shortest time exploring both the novel and familiar objects (P < 0.001). These results suggest that the existence of portosystemic collateral circulation per se may be responsible for subclinical encephalopathy.

Introduction

Hepatic encephalopathy (HE) is a neuropsychiatric syndrome observed in patients with liver failure and/or portal-systemic bypass. It is frequently considered to be a complex syndrome involving several behavioral manifestations such as personality changes, memory disorders, disorientation, flapping tremor, shortened attention span, lack of muscle coordination, bradykinesia, somnolence, and changes in sleep pattern (1). However, inconsistent results regarding the behavioral, metabolic and neurochemical characteristics indicate that the pathogenesis and pathophysiology of the syndrome are still obscure (2). Many investigators have reported that this could be caused by the lack of well-
standardized animal models (3). Although the most extensively used experimental model of HE is the end-to-side portacaval anastomosis (4), new surgical techniques have been developed to overcome the reported limitations of this model (5). Therefore, we have selected two different experimental models which simulate several main pathological features that can cause HE associated with cirrhosis: portal hypertension (PHT) and cholestasis (CHO). Both models present two frequently associated main pathologic findings of chronic liver diseases, i.e., PHT and HE (6). PHT may be an early manifestation of cirrhosis. It is defined as an increase in hydrostatic pressure of the portal vein, commonly caused by a fibrotic obstruction of hepatic blood flow (7).

Although the pathophysiological consequences of PHT per se are still not well understood, a few behavioral manifestations have been reported, such as motor incoordination (8) and marked general intellectual impairment but preserved verbal ability (9). On the other hand, extrahepatic CHO, which consists of bile stagnation in the liver, involves a combination of hepatic insufficiency and PHT. Its neurobehavioral outcomes have been partially investigated and some authors have reported clear neuropsychological impairments like fatigue and memory loss (10), together with anatomical changes in the central nervous system (CNS) (11). However, the behavioral impairments occurring in the two experimental models of PHT have not been compared. Thus, in the present study, we tested working memory ability in CHO and PHT rats using an original behavioral procedure that is highly sensitive to impairments of the hippocampal formation, a CNS structure directly related to learning and memory function in rats.

**Material and Methods**

Male Wistar rats (225-250 g) were housed in cages in temperature-controlled rooms and received water and food *ad libitum*. All animal treatments and surgical procedures employed in this study were performed in accordance with the principles and practices of the 1986 Guidelines for the Care and Use of Laboratory Animals (EC 86/609), published in Spain in RD 223/1988. Animals were randomly divided into three groups: a control group (N = 20), a group of rats with microsurgical CHO (N = 20), which underwent a resection of the bile ducts that drain each hepatic lobe in continuity with the common bile duct (12). These rats showed increased hepatobiliary parameters and histological lesions which confirm CHO. And a third group of rats with extrahepatic PHT (N = 20) induced by a triple portal vein ligation which causes a calibrated vascular stenosis (13) as assessed by the measurements of PHT.

The behavioral test was carried out 60 days after surgery in all groups. A spontaneous object recognition memory test was applied using a constantly illuminated wood open-field arena (dimensions: 65 x 45 x 45 cm). The set of objects selected for visual discrimination was made of wood and plastic and objects were available in triplicate. The texture and shape of the objects were similar to those described by Ennaceur and Delacour (14). Initially, the animals were placed inside the box and were allowed to explore it for 3 min during habituation. Thereafter, each rat was tested in two different sessions separated by a 48-h interval. Each session consisted of two different trials: in the first trial (T1 or sample phase) two identical objects were placed in the corners of the box. In the second trial (T2 or test phase) the objects were replaced with a new set (novel and familiar objects). The position of the objects was randomly changed for each animal and session. The time allowed for exploration in each trial was 3 min. The following variables were analyzed according to the procedure proposed by Ennaceur and Delacour (14): 1) Discrimination index: the
difference between the time spent to explore the new object (B) and the time spent to explore the familiar one (A) during the test phase (T2). 2) Index of habituation to the exploratory behavior defined as the total exploration time of the objects in the sample phase (T1) and test phase (T2). 3) Index of habituation to the sample: time spent to explore the familiar object (A) in the sample phase (T1) and the test phase (T2).

Data were analyzed statistically by one-way analysis of variance (ANOVA) with time of exploration as the measured variable and the experimental group as independent variable. *Post hoc* Newman-Keuls tests were applied. The results were considered to be significant when P < 0.05.

**Results**

There were significant group differences in the following behavioral indexes: discrimination between objects (F2,57 = 26.5, P < 0.001) and habituation to the sample (F2,57 = 17.6, P < 0.001). Further analysis revealed that the control group was able to discriminate between the novel and familiar objects during the test phase compared to the CHO and PHT groups (Figure 1). Although no statistically significant differences were found between the control and CHO groups in the time spent exploring the familiar object, a significant decrease in the time spent exploring the novel object was found in the CHO group compared to the control group. On the other hand, no significant differences were found in the three groups regarding total time used for exploration during the sample and test phases (F2,57 = 0.618, P = 0.543), as shown in Figure 2. Moreover, only control rats showed habituation to the familiar object during the test phase (Figure 3). Finally, the PHT group showed the poorest performance in all of the behavioral indexes measured, as evidenced by the limited time spent exploring the available objects.
Discussion

Our results indicate that, unlike the control group, the CHO and PHT groups showed an impaired ability to discriminate between the novel object and the previously experienced sample object. In the literature, this novel object preference is termed “object recognition”. However, a more serious memory impairment was observed in rats with CHO because they used excessive time to explore the familiar and novel objects, as shown in Figure 3. The animals with PHT, in turn, also displayed preference for the novel object although their exploration time was shorter than that spent by the control rats (Figure 3). The clear memory impairment of the CHO group indicates that PHT alone is not a sufficient condition to give rise to some of the main characteristics of HE.

The microsurgical extrahepatic CHO technique prevents both the formation of hilar biliary pseudocysts and of abdominal-thoracic abscesses, complications which are common in macrosurgical CHO (15,16) and, therefore, reduces mortality compared to the bile duct ligation model (17). Since this microsurgical CHO model does not evolve with infection, it would be more appropriate than the macrosurgical technique of common bile duct ligation and section to study hepatic insufficiency and its consequences, among them, HE.

Thus, endotoxemia may be involved in the HE associated with CHO in addition to other factors such as hyperammonemia. Liver endotoxemia can stimulate Kupffer’s cells to produce inflammatory cytokines such as tumor necrosis factor-α, interleukin-6, and interleukin-1β. These pro-inflammatory cytokines produced at the periphery can function as hormones and, since they can act in the CNS, they could produce HE.

It is clear from the results obtained in the present study that CHO-induced encephalopathy represents a useful experimental model to simulate some of the neuropsychological consequences of HE. In the first place, a similar impairment in working memory has been reported in patients with HE (18). In the second place, the time interval chosen (60 min) seems to involve the activation of hippocampal formation and not the prefrontal cortex (19). The hippocampus has been extensively related to recognition memory in rats (20) or its equivalent in humans (21,22). Recent data support the view that a reduction in long-term potentiation, the neural mechanism of memory found mainly in the hippocampus, could be responsible for the intellectual impairment reported in HE (23). This mechanism would favor the hyperammonemia hypothesis of HE since impairments in conditional discrimination behavior have also been reported in an animal model of hyperammonemia (24). Therefore, the working memory impairment induced by CHO could be caused indirectly by hyperammonemia, a common consequence of CHO (25). Although the molecular mechanisms of the role of ammonia in the neurological damage found in HE are still being discussed, the present study provides support for the ammonia hypothesis of HE and validates the CHO animal model of HE.

The etiology of HE arises from the gut, where toxic substances derived from food, drugs and bacterial metabolism are not detoxified by the liver, but are directly exposed to the brain causing cerebral disturbances. The etiology of HE, therefore, is probably multifactorial, with elevated blood ammonia levels playing an important and central role (26).

One of the mechanisms of ammonia detoxification in the liver is the urea cycle. The brain does not have a urea cycle and metabolizes ammonia to glutamine, causing a reduction in the intracerebral levels of glutamate, which is one of the most important excitatory neurotransmitters. The ammonia toxicity is probably more complex because, among other things, it also stimulates nitric oxide synthase causing cere-
brovascular disorders, and benzodiazepine receptors, which stimulate the GABA receptor (the most important neurotransmission inhibiting receptor) by generating neurosteroids (26). Current evidence suggests that neurotransmission changes rather than brain energy failure are the primary cause of HE. Liver failure results in altered expression of several genes coding for proteins having key roles in the control of neuronal excitability. Such alterations include decreased expression of the glutamate transporter-1, and increased expression of monoamine oxidase-A isoform, the “peripheral-type” benzodiazepine receptor, as well as of constitutive neuronal nitric oxide synthase. These changes result in altered protein expression and in increased extracellular brain glutamate, increased degradation of monoamine neurotransmitters, increased synthesis of neurosteroids with inhibitory properties, and increased production of nitric oxide in the brain in chronic liver disease (27,28). Prehepatic PHT by partial portal vein ligation leads to morphologic changes in the rat’s CNS, including alterations of the blood brain barrier, astrogliosis and angiogenesis in the hippocampus, as well as functional changes in the integrity of the blood brain barrier and subtle impairments in brain activity which manifest as behavioral alterations such as motor incoordination. Thus, in the present study, we observed that the time of object exploration was reduced in rats with PHT. However, these rats tended to interact more with a familiar object in a free-choice situation, suggesting the development of a subclinical form of HE (29).

The experimental models of microsurgical CHO and PHT share intestinal bacterial translocation, development of portosystemic collateral circulation, and increased hypothalamic levels of leu-enkephalin (30), although these alterations are more intense in CHO. This could be the reason why HE, assessed in this study by memory impairment, was also more intense in CHO while it was occasionally subclinical in rats with PHT, producing a mild alteration in the object recognition task.

A new nomenclature of HE has been used since 2002, classifying HE into encephalopathy type A, associated with acute liver failure, type B, associated with portal systemic bypass without intrinsic hepatocellular disease, and type C, associated with liver cirrhosis (31). According to this new classification, CHO-induced HE would correspond to type C whereas PHT, in which there is only portal-systemic circulation without hepatic insufficiency, would occur with subclinical HE named type B in the human clinical area.

References


