Spatial memory deficits in juvenile rats with pilocarpine induced temporal lobe epilepsy

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One of the most frequent forms of epilepsy in humans is temporal lobe epilepsy. Characteristic to this form of the disease is the frequent pharmacoresistance and the association with behavioural disorders and cognitive impairment. The objective of our study was to establish the degree of cognitive impairment in a rat model of temporal lobe epilepsy after an initial epileptogenic exposure but before of the onset of the effect of long-duration epilepsy.

Methods. For the experiment we used 11 rats. Status epilepticus was induced by systemic administration of a single dose of pilocarpine. The animals were continuously video-monitored to observe the occurrence of spontaneous recurrent seizures; during weeks 9-10 we performed eight-arm radial maze testing in order to assess the cognitive impairment.

Results. Animals developed spontaneous recurrent seizures after a 14-21 day latency with a daily average seizure density of 0.79±0.43 during weeks 9-10. Epileptic rats had significantly more working memory errors per session, more reference memory errors and the number of visited arms was also significantly higher. Accuracy was also lower in the pilocarpine treated group. Interestingly significant differences disappeared after six days of trials.

Conclusions. Our study shows behavioural deficits occurring after 9-10 weeks of epilepsy in the pilocarpine model of epilepsy applied to juvenile rats. In contrast to previous studies, we showed that juvenile rats with short duration of epilepsy are able to learn the behavioural task, therefore a morphopathological and/or behavioural "no-return point" regarding the development of severe cognitive impairment is not reached by status epilepticus alone.

Keywords: epilepsy, hippocampus, cognitive impairment.

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Introduction

Epilepsy defines a spectrum of disorders that are characterized by the unpredictable occurrence of seizures, widely differing seizure types and causes, and varying impacts on individuals and their families. The disease creates a significant socio-economic burden not only at the individual and family level, but at the societal level, due to the implications regarding health services. Currently there is no cure available, and all therapeutic choices are only symptomatic, with the goal to eliminate or at least to alleviate seizures without interfering with normal function. Data from recent years suggests that by accounting only for seizures, we grossly underestimate the true impact of epilepsy, as the disease is frequently associated with comorbid cognitive and behavioural impairment.

One of the most common forms of epilepsy in humans is temporal lobe epilepsy (TLE), which is also the type (1) most prone to pharmacoresistance and (2) more frequently associated with behavioural disorders and cognitive impairment. TLE causes deficits in declarative memory and in the performance of visuospatial tasks, reduction in psychomotor speed, and executive functions and, on a long term, cognitive deterioration. As such, psychiatric disorders and cognitive impairment in patients with epilepsy may be considered to reflect the degree of brain dysfunction.

One particularly interesting brain region is the hippocampus which is implicated in memory and learning as well as pathological synchronous oscillations (i.e. epileptic seizures). Therefore the hippocampus may be involved in the same time in temporal lobe epilepsy and the cognitive impairment associated with the disorder.

The cognitive deterioration during chronic epilepsy was described earlier, but there is a lack of data regarding the exact time-course of the development of memory impairment. Therefore, we studied cognitive impairment in a rat model of epilepsy after an initial epileptogenic exposure but before the onset of the effect of long-duration, chronic epilepsy.

Methods

Animals

For the experiments we used 11 male Wistar rats (100-250g) that were housed under standard conditions: constant temperature (21-24°C) and humidity, 12 hours light/dark cycle and access to food and water ad libitum. All procedures involving animals were carried out in accordance with EU Directive 2010/63/EU and national and local guidelines and policies (CEC 26/2011). Five 5-6 week old rats were used for status epilepticus induction as well
as sixe age- and batch matched control animals. Rats were randomly assigned to either control or pilocarpine treated group.

Pilocarpine protocol
The pilocarpine model of epilepsy was described earlier: rats receiving a single dose of systemic pilocarpine develop status epilepticus (SE); after a period of latency, spontaneous recurrent seizures (SRS) occur that gradually will become more frequent and severe, leading to a form of epilepsy which is analogous to chronic epilepsy in human patients. Five rats received a single injection of methylscopolamine (1 mg/kg, s.c., necessary to counteract the peripheral effects of the cholinergic agonist) 30 minutes prior to pilocarpine (350 mg/kg, i.p.) injection. The animals behaviour was observed by experimented researchers for at least 5 hours afterwards. Within the first 10-30 minutes after pilocarpine injection all rats developed seizures with variable severity that evolved into status epilepticus. Seizures were interrupted 2 hours later by administration of Diazepam (5 mg/kg, i.p.). The duration of SE was selected in accordance with literature data in order to reduce mortality, but in the same time to allow for neural changes characteristic to the model to be established.

Continuous video monitoring
All rats were continuously video monitored during and after SE for a period of 120-150 days. Night-vision motion-capture camera with i-Catcher software (iCode Systems Ltd, Hampshire, England) was used for video-recording. After the appearance of SRS we established the early seizure pattern and classified seizures according to the modified Racine scale. Video recording were analyzed by several researchers, but, in order to exclude inter-observer differences, a single person reanalyzed all putative SRS found.

Eight-arm radial maze testing
Rats were tested in the eight-arm radial maze (RAM) according to the procedure described by Olton. First, all animals (control and pilocarpine treated) were partially food deprived until they reached 85% of their body weight. For testing we used a handmade wooden brown eight-arm radial maze with walls but without plastic covering, constructed according to standard specifications. Each of the arms projected from one side of an octagonal center. One small fragment of food pellet was placed at the far end of 6 of the 8 arms before each trial. The same arms remained unbaited throughout the protocol. We did not use reinforcement (i.e. replacement of the acquired pellets). During each trial individual animals were placed in the octagonal center and left in the maze until they had recovered all 6 food baits or until 10 minutes had elapsed, whichever occurred first. Placing all four paws inside an arm was considered entrance. The trial was repeated at least six consecutive days (one test per day), always within the same time interval and same conditions (including visuospatial clues); times of arm entrances, the identity of each arm entered, the order of arms, the number of baits recovered as well as the time to finish the task was recorded. We analyzed the following parameters: working memory error (WM) – defined as return to a previously visited arm during the same trial, reference memory error (RM) – entry to an unbaited arm after day 2, accuracy of choice (ACC) – number of obtained food pellets from the first 6 visited arms, total number of visited arms (TO) and the number of food pellets obtained during each completed trial (FO). Day one was excluded from analysis (adaptation to the new environment).

Statistical analysis
Descriptive statistics, nonparametric tests (two-tailed Mann-Whitney) and general linear model were applied for data analysis using GraphPad Prism 5.00 for Windows (GraphPad Software, Inc., San Diego, California, USA). Unless otherwise stated all values are given as Mean±SE. A p value less then 0.05 was considered significant.

Results

Seizure pattern
Rats that underwent pilocarpine induction developed SRS after a latency period of 14-21 days. We considered for analysis only rats that developed Racine stage 5 seizures (which is homologous to secondary generalized seizures in humans). The average duration of SRS was 41.9±15.4 second (Mean±SD). The number of seizures per day reached 0.79±0.43 during weeks 9-10 (all time periods are relative to time of induction with pilocarpine), the period of RAM testing. It must be noted however that the seizure pattern’s further evolution was also monitored (i.e. after the RAM testing) to prove the development of chronic epilepsy (Figure 1).

Fig. 1. Seizure pattern progression in rats with induced TLE.
Effects of the pilocarpine induced TLE in juvenile rats on the behaviour in the eight-arm radial maze

Rats from the control group learned the task much quicker than rats with TLE as the working and reference memory errors were low starting from day 2 and they further diminished during the 6 day trial period (for summary of results see Figure 2). Furthermore, accuracy was high from the beginning and the rats in this group obtained all food pellets starting from day 2 (Figure 3).

Conversely, rats treated with pilocarpine (i.e. rats with TLE) had significantly more working memory errors (6.4±0.8 versus 2.3±0.3, p=0.0026), more reference memory errors (2.86±0.25 vs. 1.88±0.22, p=0.023) and the number of visited arms was also significantly higher (13.3±0.76 vs. 9.7±0.4, p=0.008). Accuracy was lower in the pilocarpine treated group (3.2±0.39 vs. 4.23±0.17, p=0.04) and it took 6 days for rats to be able to collect all food pellets. It must be noted however, that significant differences appeared only during the first days. Most interestingly epileptic rats eventually were able to learn the task, albeit delayed, and on day 6 no significant changes could be found, regardless the analyzed parameter (Figure 2).

Discussion

Although several papers present the behavioural aspects of TLE in different conditions – immature and adult rats, after a short period or longer period of seizures and with
different induction age – there is a lack of data regarding the behavioural consequences of SRS in epilepsy induced in juvenile rats (postnatal 4-5 weeks).

All previous studies suggested that spatial learning in epileptic rats is severely impaired. Interestingly, our data suggests that, although learning the task in the eight-arm radial maze is difficult, it is not impossible for juvenile epileptic rats. In our study we observed a gradual improvement of the performance of the epileptic rats along the 6 days of testing. This improvement was characteristic both to the reference and working memory; also the gradual reduction of the number of arms visited showed a learning and goal oriented behaviour, that eventually lead to the collection of all food baits.

In contrast to our study, several groups found that inducing epilepsy in immature rats – postnatal 14-17 days, roughly the equivalent to 6-12 months of human development – causes a permanent reduction in the performance in RAM. This effect however may be caused in conjunction with permanent changes in motor performances. There are morphological, behavioural and psychological evidences that show a link between early cellular alterations and long-term impairment of hippocampal function in adulthood. Furthermore, the development of the dentate gyrus (DG), which plays a significant role in learning, is mainly completed during the first two weeks; as such, it is possible that in previous studies the observed long-term effect on spatial memory is due to the result of seizures in the developing circuitry of the DG.

Other studies looked at the effect of SRS over a long period of time and found that after 5-months duration of epilepsy, rats are unable to learn the task in the RAM, i.e. spatial memory and learning capabilities is adversely affected. The combination of lesions in the hippocampus and in layers II and III of the entorhinal cortex (EC) causes impairment of learning and retention, therefore supporting the poor performance of rats in RAM testing. Furthermore, longer period of SRS significantly worsens the neuronal loss both in the hippocampus and the EC. Therefore, it seems that both SE and SRS contribute to the neuronal loss and the subsequent behavioural impairment in chronic epilepsy.

When shorter periods of SRS were investigated, only in adult animals, severe memory impairment was found with a permanent learning disability. In contrast to this finding, our animals were able to eventually learn the task in RAM, and as the duration of epilepsy was quite short, it seems that the initial trigger (i.e. SE) alone is not enough to disrupt the neuronal network and cause permanent learning disability, at least in juvenile rats. Further studies, morphopathological and chronic performance testing of juvenile rats, are needed to elucidate the onset of “no-return point” in the development of severe cognitive impairment in epilepsy.

Conclusion
In conclusion, the present study shows behavioural deficits occurring at 9-10 weeks after induction of epilepsy with pilocarpine in juvenile rats. In contrast to previous studies, we showed that juvenile rats with short-duration epilepsy are able to learn the task in RAM, therefore a morpho-pathological and/or behavioural “no-return point” regarding the development of permanent learning disability and severe cognitive impairment is not reached by the initial SE alone. Our study also confirms that the pilocarpine model of epilepsy combined with RAM testing is a useful tool to explore behavioural disorders in general and cognitive decline in particular in rats.

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