Punicalagin effect on total sleep deprivation memory deficit in male Wistar rats

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Sleep deprivation has deteriorating effects on cognitive functions and activation of brain inflammation mechanisms has been reported by some studies following total sleep deprivation. Some studies have reported the health benefits of punicalagin, a main abstract from Punica granatum L., including those for the treatment of Alzheimer’s disease. The antioxidant characteristic of punicalagin and the fact that sleep deprivation accelerates mediators of inflammation led us to further explore the possible neuroprotective role of punicalagin in total sleep deprivation memory impairment in a rat model. In this study, male Wistar rats were implanted with a cannula in the lateral ventricle to receive intracerebroventricular injections (drug or vehicle). The animals were trained for the passive avoidance test and then received intracerebroventricular injections of different doses of punicalagin (0.001, 0.01, or 0.1 μg/rat). Then, they were placed in the sleep deprivation apparatus for 24 hours and tested afterwards for memory retrieval and locomotion. Our results indicated that 24 hours of total sleep deprivation impaired memory processes. PG microinjection before TSD did not prevent the deteriorating effect of total sleep deprivation on memory, and only showed a tendency of restoring the memory impairment. Comparison of the locomotor activity between the animals in different groups showed a significant increase in the total sleep deprivation sham groups that received two of the highest doses of punicalagin. Considering the reported beneficial actions of PG by other studies, further investigation is needed into the possible effects of PG in memory alterations.

Keywords
Punicalagin (PG); Total sleep deprivation (TSD); Memory deficit; Male rats

1. Introduction

Sleep deprivation has profound adverse effects on cognitive functions including information processing speed [1] and learning and memory [2, 3]. Sleep deprivation not only leads to a decline in cognitive functions but also generates higher levels of inflammatory cytokines [4, 5]. Previous studies have indicated that physiological and biochemical morbidity following sleep disorders is related to oxidative stress [6–8]. Besides, cognitive decline following sleep deprivation has been reported to correlate with increased oxidative stress in the brain and body [9, 10].

The cycle of sleep, as a homeostatic process, is generally divided into two main phases: rapid eye movement (REM) and non-rapid eye movement (NREM) [11]. Theta waves are dominant in REM sleep and the phase is accompanied by low muscle tone. On the other hand, EEG in NREM sleep is composed of waves with high amplitude and low frequency, and the phase is accompanied by decreased muscle activity [12, 13]. Sleep cycle in rats takes about 12-20 minutes [14] and the NREM sleep-related model of REM-sleep proposes that REM sleep is homeostatically linked to NREM sleep rather than to waking and occurs in response to NREM-sleep expression to compensate certain processes of NREM sleep [15, 16]. Similar to the sleep cycles, sleep deprivation is divided into total sleep deprivation (TSD) and rapid eye movement sleep deprivation (RSD). Reports showed that TSD impaired consolidation of declarative memory [17] and motor adaptive task memory [18]. However, selective REM sleep deprivation disrupted spatial memory consolidation in an eight-box task [19] and water maze [20]. TSD and RSD both inhibit the induction of long-term potentiation (LTP) [21, 22]. Considering that REM sleep is not the only phase that is involved in the consolidation of memory [23], in the present study we selected a 24-hour TSD model to investigate its effects on memory deficit.

Punicalagin (PG), as a potent antioxidant, is a hydrolyzable polyphenol in Punica granatum L. and is a major component of pomegranate responsible for its health benefits [24]. PG also has an anti-amyloid beta (1-42) fibril aggregation effect, a potency that can be considered in the treatment of Alzheimer’s disease [25]. The usefulness of pomegranates has also been reported by a clinical study for memory recovery after ischemic stroke [26]. Nevertheless, several issues central to a basic understanding of the modulation of cognitive performance and the possible prevention of memory decline by this natural herbal component remain unresolved.
The mentioned antioxidant characteristic of PG and the fact that sleep deprivation accelerates mediators of inflammation inspired us to investigate the possible neuroprotective effect of PG in a TSD memory impairment model in rats.

2. Materials and methods

2.1 Animals

Male Wistar rats (weighing 200-220 g, age 9-11 weeks) were purchased from Institute for Cognitive Science Studies and kept in standard conditions (Temperature: 23 ± 1 °C; light-dark cycle: 12:12 h; lights on at 7:00 p.m.) in an animal house for one week before the experiments. The animals had free access to water and food and kept in groups of 4 in Plexiglas cages. Each experimental group included 8 rats and each animal was used only once. All the experiments were conducted between 9:00 a.m. to 1:00 p.m. under the Guide for the Care and Use of Laboratory Animals (National Institutes of Health Publication No. 80-23, revised 1996). The experiments were approved by the Research and Ethics Committee of Tehran Medical Sciences, Islamic Azad University, Tehran, Iran.

2.2 Drugs

Punicalagin was purchased from Sigma-Aldrich (≥ 98% (HPLC); Cat. No.: P0023) and based on pilot studies doses of 0.001, 0.01, and 0.1 µg/rat were prepared by first dissolving PG in 0.1% alcohol (Merck, Germany, 99%) and then diluting the solution with normal saline (0.9%). All doses of PG were prepared freshly before injection. Ketamine (Alfasan, Holland) and Xylazine (Pantex, Holland BV) were purchased and used for stereotaxic surgery.

2.3 Study design

Fig. 1 illustrates the general design of the study. The explanation for each step has been provided in the respective following sections:

2.3.1 Stereotaxic surgery

Each rat was anesthetized using 2 mL/kg intraperitoneal injection of ketamine hydrochloride 10% (50 mg/kg) and xylazine 2% (4 mg/kg). The animal’s head was then fixed in a Kopf stereotaxic frame. A stainless steel guide cannula (22 gauge) was implanted above the left lateral ventricle (AP = -0.9, ML = 1.3, DV = -4.5 from skull surface) according to rat brain atlas by Paxinos and Watson [27] and fixed to the skull with instant glue and acrylic resin. After surgery, an injection of meloxicam (1 mg/kg; s.c.), as a painkiller, was made and followed by an injection of penicillin-G 200,000 IU/mL (0.2-0.3 mL/rat, single dose, intramuscular). A stainless-steel styllet was inserted into the guide cannula to prevent its occlusion during the recovery period (one week).

2.3.2 Intracerebroventricular injections

To inject PG, we used a 2 µL Hamilton syringe connected by a polyethylene tube (PE 20) to an internal cannula (27-gauge, terminating 1.3 mm below the tip of the guide cannula). Each injection (2 µL) was made over a 2 min period and the injection cannula was left in place for an additional 30 sec to prevent the backflow of fluid. Injections were made following the training section of the passive avoidance test and the animal was placed in the sleep deprivation apparatus after 30 minutes. The animals could not sleep during the 30-min period due to being handled by the experimenter.
2.3.4 Passive avoidance memory test

Passive avoidance emotional memory test apparatus included a Plexiglas box composed of two equal-sized partitions (20 × 20 × 30 cm³) separated by a wall, in which a guillotine door was installed to permit rat’s passage when necessary. The white partition was lit with a 25-watt electric lamp hanged 50 cm above the apparatus floor. The floor of the dark partition was made of steel rods (1 cm intervals from each other) connected to a stimulator to provide foot shocks (Settings used: Squarewave, frequency 50 Hz, 1 mA applied for 3 seconds) generated by an insulated stimulator [28].

2.3.4.1 Habituation, training, and retrieval test. To habituate the animals to the passive avoidance apparatus, the animals were allowed to habituate in the experiment room for at least 30 min prior to the experiments. Each animal in each experimental group was gently placed in the white partition of the apparatus and then, the guillotine door was raised 5 sec later. As rats are inclined to move to the dark partition, the entrance latency to the dark partition (step-through latency; STL) was recorded when the rat placed all four paws in the dark partition. The gate was closed after 10 sec and the animal was returned from the dark partition into the home cage. Animals with a STL more than 100 sec were excluded from the experiments (2 rats).

The training trial was performed 30 min after the habituation. The animal was placed in the light compartment and 5 sec later, the guillotine door was opened. As soon as the animal crossed to the dark compartment, the door was closed and a foot shock was immediately delivered to the grid-floor of the dark compartment. After 20 sec, the animal was removed from the apparatus and temporarily placed in its home-cage. The procedure was repeated in 2 min time. The training was terminated when the rat remained in the light partition consecutively for 120 sec. The number of trials (entries into the dark partition) was also recorded. All animals learned within a maximum of three trials.

For the retrieval test of the long-term memory, each animal was placed in the light partition 24 hours after the training for 20 sec, then the door was raised and the STL to enter the dark partition was measured. The test session was ended once the animal entered the dark partition. A cut-off time of 300 sec was applied for those animals which remained in the light partition. No electric shock was applied during the retrieval session [28, 29].

2.3.5 Measurement of locomotor activity

Following the passive avoidance test, the rats were tested using a locomotor activity test apparatus (BorjSanatazma Co, Tehran, Iran). The animals were acclimatized to the test room (15 min) and then put into a clear Plexiglas container box as the test chamber (30 × 30 × 40 cm³). The apparatus has a gray Perspex panel (30 × 30 × 2.2 cm³) with 16 photocells dividing the box into 16 equal-sized squares. The number of crossings from one square to another during a 5-min period was measured as the locomotion index [30].

2.4 Histological verification

To verify the cannula placement, each rat was deeply anesthetized with Ketamine and Xylazine, the brain was removed and kept in formaldehyde (10%) for 5 days, the brain was sliced (50 µm) and the cannula path into the left ventricle was verified. The data of misplaced injection sites were eliminated (Three animals).

2.5 Experimental design

Seven groups of animals were used in this study (Table 1). The animals that received surgery were given one week recovery period and then the passive avoidance training was conducted. Vehicle or PG injection was then performed and the animals were put in the TSD apparatus. The apparatus was either kept off or turned on for the sham and experimental groups, respectively. Then, all the animals underwent the passive avoidance test followed by the locomotion test.

2.6 Statistical analysis

Data were analyzed by GraphPad Prism® (Version 9.0). Kolmogorov-Smirnov statistical test did not show the normality of STL data for all the experimental groups. Thus, a Kruskal-Wallis test followed by Dunn’s multiple comparisons was used to identify possible differences. STL data are expressed as median (interquartile) values. The data regarding distance traveled followed a normal distribution in all groups and were analyzed using one-way ANOVA test followed by Tukey-Kramer post-hoc test, as appropriated. Distance traveled data are expressed as mean ± SEM (standard error of the mean). P-values less than 0.05 were considered to be statistically significant.
3. Results

3.1 Effects of TSD on the retrieval of long-term memory

To test the possible effects of TSD on the retrieval of long-term memory, Kruskal-Wallis test was used on control groups data (normal control, TSD sham control, and TSD control). The test revealed that there was a statistically significant difference between all the three groups being tested [Chi-Square (2) = 11.54; P < 0.01]. Post-hoc analysis with Dunn’s multiple comparisons revealed a significant decrease in median STL of the animals that received 24 hours of TSD (TSD control group) as compared to the normal control group (P < 0.01; Fig. 2A).

To analyze locomotor activity, univariant one-way ANOVA test was performed. The test revealed no statistically significant difference between all the three groups being tested [F (2, 21) = 1.820; P = 0.1867; Fig. 2B].

Our data revealed that 24 hours of TSD impaired retrieval of long-term memory without affecting the locomotor activity.

3.2 Effects of intracerebroventricular microinjection of PG on the retrieval of long-term memory following TSD

Comparison of the TSD sham groups using Kruskal-Wallis test revealed that there were no statistically significant differences between the groups being tested as compared to the vehicle group [Chi-Square (3) = 6.001; P = 0.1852; Fig. 2A-TSD sham groups]. Comparison of the TSD groups using Kruskal-Wallis test also revealed no statistically significant differences between the groups being tested, compared to the vehicle group [Chi-Square (3) = 7.68; P = 0.0471; Fig. 2A-TSD groups].

Comparison of the TSD sham groups using one-way ANOVA on the results obtained from locomotion activity tests indicated a statistically significant difference between the experimental groups being tested [F (4, 35) = 15.20, P < 0.0001; Fig. 2B-TSD sham groups]. Post-hoc analysis with Tukey-Kramer test revealed that the two highest doses of PG (0.01 and 0.1 μg/rat; P < 0.0001 and P < 0.0001) had significant effects on the locomotion (Fig. 2B-TSD sham groups). Application of one-way ANOVA on the results obtained from locomotion activity tests of the TSD groups indicated a statistically significant difference between all the experimental groups being tested [F (4, 35) = 1.259, P = 0.3047; Fig. 2B-TSD groups].

Our data revealed that none of PG doses altered memory consolidation by themselves (TSD sham groups). Also, PG microinjection before TSD did not alter memory consolidation significantly and only a tendency of memory restoration was observed. While PG changed the locomotive activity in the TSD sham group, no changes was observed in the TSD group.

4. Discussion

Lifestyle changes in modern societies have made sleep deprivation a major challenge that can increase the risk of some neurologic diseases [31]. Sleep plays a prominent role in the
maintenance of mental and physical performance and inadequate sleep has deteriorating effects on cognitive functions [32] including memory functions [33].

In line with previous studies [28, 34], we here showed that acute TSD impaired memory processes in the passive avoidance task. The data also showed that PG, per se, did not have any significant effects on the memory functions in this test, but it increased the locomotor activity at the two highest applied doses. PG microinjection before TSD only showed a tendency of restoring the memory impairment and did not alter the locomotor activity. Hence, the observed tendency was not under the influence of changes in locomotor activity. Although some reports indicated no changes in locomotor activity following TSD [35, 36], other reports have indicated an increase in locomotion following sleep deprivation [37] and a recent research on the effects of pomegranate on parkinsonism proposed changes in the level of brain-derived neurotrophic factor (BDNF) as a possible mechanism for the observed increase in the locomotor activity [38]. Applied methods for sleep deprivation and the duration of sleep deprivation might be the reason for the observed differences. Sleep deprivation leads to brain inflammation [10, 39], and reports indicate the protective effect of PG in brain inflammation [40, 41].

Our results showed that 24 hours of TSD impaired retrieval of memory in the passive avoidance test. Signaling pathways underlying the deficit in LTP have been investigated by some researchers. For example, the expression of BDNF and its downstream targets (Synapsin I, cAMP response-element-binding, CREB, and calcium-calmodulin-dependent protein kinase II, CAMKII) reduced in the hippocampus following 8 and 48 h of sleep deprivation [12]. In addition, the role of a compensatory increase in cAMP signaling [42] and reduction of extracellular adenosine [43] in preventing LTP deficit following sleep deprivation have been reported. Maintaining CREB at its beneficial level might be a possible mechanism for the observed tendency, as a study showed that hydro-alcoholic leaf extract of Terminalia cattapa, which also contains PG, regulates BDNF and CREB levels [44].

Sleep deprivation promotes the production of pro-inflammatory proteins both in humans and rodents [45, 46]. The anti-inflammatory effects of PG have been reported in rats [41] and its neuroprotective effect on glutamate-induced oxidative stress has been reported in the mouse hippocampal cell line, HT22 [47]. It has also been reported that PG can inhibit lipopolysaccharide-induced inflammation in certain types of macrophages [40] and anti-inflammatory property of PG works against memory deficit via the prevention of neuroinflammation [48]. Furthermore, it has been reported that PG inhibited the expression of NF-κB and the inflammatory proteins, expression of which is mediated by NF-κB in mice, resulting in the reduced expression level of TNF-α, IL-1β, iNOS, Cox-2, ROS, and NO in the brain [48]. Hence, another possibility that must be considered is the antioxidant and anti-inflammatory properties of PG.

It has been shown that sleep stages and its pattern vary in different animals, which is a point for consideration while the results are interpreted [49]. For example, other studies are reporting that sleep deprivation, for 96 hours, did not induce necrotic or apoptotic cell loss in rat’s brain [50]. Another study that measured antioxidant and oxidant markers in the hippocampus reported short- and long-term memory impairment after 72 hours of sleep deprivation, but found no significant differences in the markers compared to the control group [51]. Also, another study reported no significant rise in oxidative stress in 24-hour sleep-deprived rats [52].

5. Conclusions
PG only showed a tendency of restoring the memory impairment following 24 hours of TSD in the passive avoidance test. Considering the reported beneficial actions of PG by other studies, further investigation is needed into the possible effects of PG on memory alterations.

Abbreviations
ANOVA, Analysis of Variance; BDNF, Brain-derived neurotrophic factor; CAMKII, Calcium-calmodulin-dependent protein kinase II; CREB, cAMP response-element-binding; LTP, Long-term potentiation; NREM, Non-rapid eye movement; PG, Punicalagin; REM, Rapid eye movement; RSD, Rapid eye movement sleep deprivation; TSD, Total sleep deprivation.

Author contributions
Shahram Zarrabian and Mohammad Nasehi were responsible for the study concept and design. Mohammad-Hossein Mohammadi-Mahdiabadi-Hasani contributed to the acquisition of behavioral data. Shahram Zarrabian, Mohammad Nasehi, and Mohammad-Reza Zarrindast assisted with data analysis and interpretation of findings. Shahram Zarrabian drafted and finalized the manuscript. Shahram Zarrabian and Mohammad Nasehi provided critical revision of the manuscript for important intellectual content. All authors critically reviewed content and approved the final version for publication.

Ethics approval and consent to participate
All the experiments were conducted under the Guide for the Care and Use of Laboratory Animals (National Institutes of Health Publication No. 80-23, revised 1996). The experiments were approved by the Research and Ethics Committee of Tehran Medical Sciences, Islamic Azad University, Tehran, Iran.

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Supplementary material
Supplementary material associated with this article can be found, in the online version, at https://jin.imrpress.com/EN/10.31083/j.jin.2021.01.378

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