Therapeutic Brief

GHK peptide prevents sleep-deprived learning impairment in aging mice

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**Abstract**

Sleep deprivation is known to cause memory impairment and is associated with inflammation and cell damage linked to neurodegenerative diseases. GHK (glycyl-L-histidyl-L-lysine) is a naturally occurring tripeptide found in mammalian plasma. GHK has anti-inflammatory activity and can pass through the blood-brain barrier suggesting the potential to prevent neuroinflammation associated with sleep deprivation. In this study, mice were injected with 15mg/kg GHK per day for five days and sleep deprived on the last two days of treatment. Sleep-deprived mice treated with GHK did not show the acute learning impairment seen in sleep-deprived mice treated with saline. GHK prevented an increase in MCP-1 and nitrotyrosine levels in the hippocampus of sleep-deprived mice suggesting that inflammatory and reactive nitrogen/oxygen species activity could be therapeutic targets for learning impairment associated with short-term sleep deprivation.

**Keywords:** GHK peptide, sleep deprivation, learning impairment, neuroinflammation, nitrotyrosine

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Sleep deprivation is an increasing problem in modern society, with sleep duration among American adults decreasing over the past 25 years. As sleep deprivation becomes the norm of the American working world, researchers are finding that a decrease in overall sleep can cause serious health issues. Both acute and chronic habitual sleep loss is related to negative impacts on mental health, cardiometabolic conditions, and pain [1]. Insufficient sleep can cause increased cytokine secretion provoking an inflammatory response in various tissues including the brain which is associated with cognitive impairment [2].

GHK (glycyl-L-histidyl-L-lysine) is a naturally occurring human plasma copper-binding peptide known to possess wound healing, antioxidant, anti-inflammatory, and anti-aging effects [3]. Because of its antioxidant and anti-inflammatory properties, GHK is a promising peptide for use in the treatment of age-related neurodegenerative conditions. This therapeutic brief describes preliminary observations on the ability of GHK to prevent the adverse neuropathological effects of short-term sleep deprivation in aging mice. The rationale for using GHK without copper was that it is important to know if GHK can be effective without its copper complex as a therapeutic parenteral injection for this type of cognitive impairment.

CB6F1 female mice were obtained from the National Institute on Aging aged rodent colony at 15 months of age. Mice were housed in 3-4 per cage in an SPF facility at the University of Washington under a 12-hour light-dark cycle starting at 6 am. The room temperature was 25℃±4. Reverse osmosis water and irradiated food (Picolab Rodent Diet 20, 5053) were supplied. All studies were approved by the University of Washington IACUC.

Mice were started on intraperitoneal injection (ip) of copper-free GHK (Peptide Sciences, Henderson, NV ) at a dose of 7.5 mg/kg (n=8), or ip saline (n=12), twice daily for five days. On days 4 and 5 of treatment, the GHK group and 6 of the saline group were sleep deprived for 4 hours as described [4]. Mice were sleep deprived in their home cage by light cage tapping and gentle stroking of the back using a small paintbrush. Six of the saline group were not sleep deprived. On the fifth day, following sleep deprivation, each group was tested in a box maze spatial navigation learning task [5]. Each mouse was given 120 seconds to find an escape hole and tested continuously for 4 trials with the escape time for each trial recorded.

Mice were euthanized by CO2 and brain fixed in formalin. Sagittal 4 µm paraffin sections of the brain were mounted on slides, which were rehydrated and incubated in a citrate buffer at 95℃ for 30 minutes for antigen retrieval. Slides were then stained for MCP-1 and nitrotyrosine to assess inflammation and oxidative stress, respectively, using specific antibodies and an HRP-DAB cell and tissue staining kit (R&D Systems). Digital images of CA3 and dentate gyrus (DG) sections of the hippocampus were taken at 20x magnifications for stain intensity analysis and processed through ImageJ with the IHC toolbox and IHC profile plugin [6].

Mice treated with GHK showed little learning impairment after sleep deprivation in line with saline-treated mice that were not sleep deprived, and significantly less than sleep-deprived mice treated with saline (Figure 1). The learning ability in the Box maze was calculated as the slope of escape times of each mouse. More negative numbers indicated faster escape times and less learning impairment. The negative slope value of sleep-deprived mice treated with GHK (SD+GHK) was similar to the negative slope value of non-sleep-deprived mice treated with saline (Control). In contrast, sleep-deprived mice treated with saline (SD) had a significant decrease in the learning curve slope indicating learning impairment without GHK.

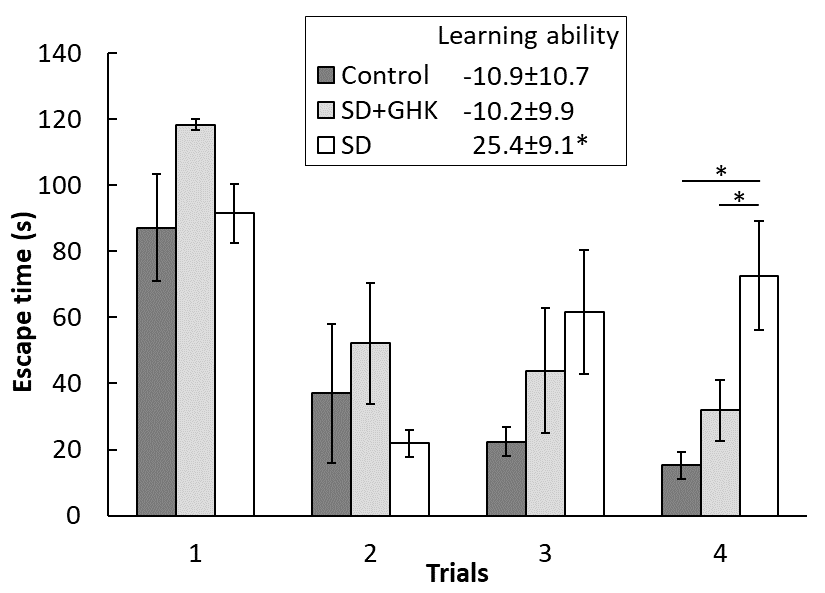


Figure 1. GHK treatment prevented learning impairment caused by sleep deprivation. The SD+GHK mouse group had escape times and a learning curve slope similar to the nonSD+saline (control) group but significantly less than the sleep-deprived saline-treated mice (SD) \*: p<0.05. Data from the nonSD+GHK cohort are not shown because there was no effect.

Inflammation and oxidative stress in sleep-deprived mice were measured by MCP-1 and nitrotyrosine staining, respectively, and quantitated by ImageJ digital analysis. MCP-1 levels in sleep-deprived mice treated with GHK were significantly lower in the CA3 region of the hippocampus compared to sleep-deprived mice treated with saline, indicating GHK's anti-inflammatory effect (Figure 2A). Notably, sleep-deprived mice treated with GHK had lower inflammation compared to both sleep-deprived and control mice in the CA3 region of the hippocampus. Sleep deprivation increased nitrotyrosine levels in both the CA3 and DG regions of the hippocampus, and GHK treatment successfully prevented such an effect (Figure 2B). GHK is known to work as an anti-inflammatory and antioxidant agent by decreasing inflammatory cytokines and reducing reactive oxygen species (ROS) levels [7]. Nitrotyrosine is considered a biomarker for oxidative stress as the result of the nitration of protein-bound and free tyrosine residues by reactive peroxy nitrate molecules formed when nitric oxide reacts with superoxide [8].

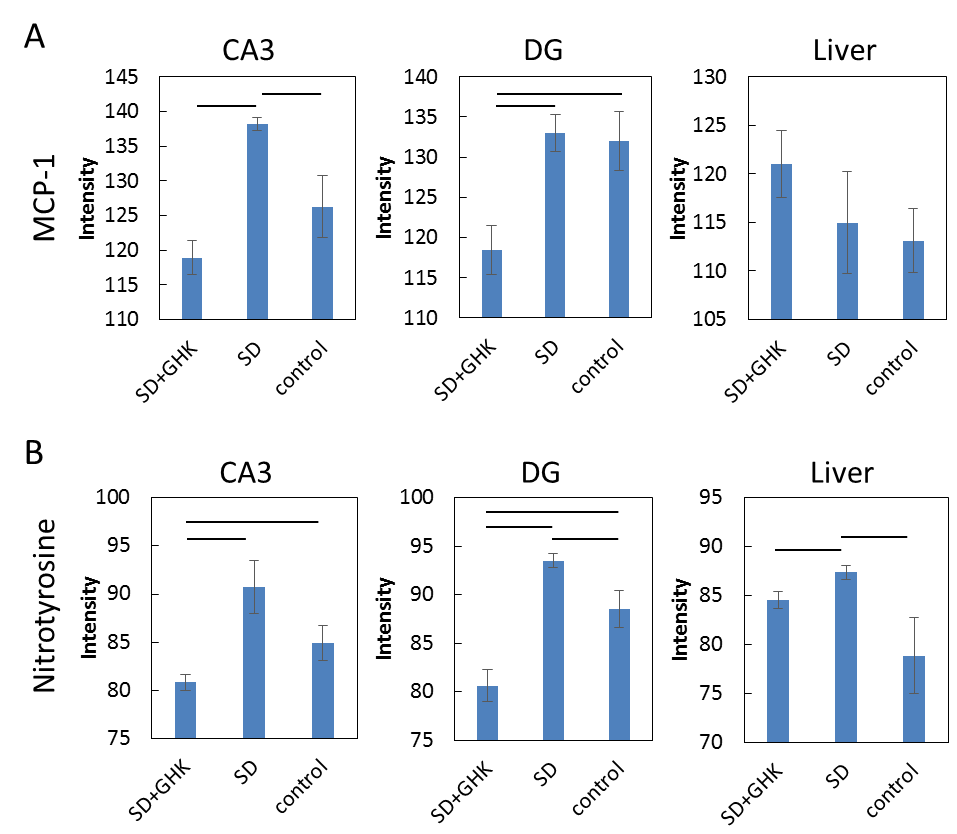


Figure 2. Immunohistochemistry images were quantified by staining intensity using ImageJ. Sleep-deprived mice treated with GHK had (A) decreased MCP-1 and (B) decreased nitrotyrosine staining intensity in the hippocampus. Results connected by horizontal lines are significantly different (*p<0.05*). Control = nonSD+saline

These results indicate that GHK can prevent sleep-deprived learning impairment associated with suppression of increased inflammation and nitrotyrosine production in the hippocampus, suggesting that GHK could be further studied as a way to prevent adverse neuropathological effects of acute sleep disruption.

**Declarations**

Authors’ contributions

All authors made contributions to the generation of data and/or writing the manuscript.

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Conflicts of interest

Warren Ladiges is a member of the Editorial Board of*Aging Pathobiology and Therapeutics*. The author declares that there are no conflicts.

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[1-7]

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