Resilience to acute sleep deprivation is associated with attenuation of hippocampal mediated learning impairment

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Abstract

Background: Sleep deprivation is a universal issue that affects individuals in different ways. While some individuals experience a deficit in performance, others experience resiliency as they maintain high levels of physical and mental activity. Sleep loss is known to cause cognitive dysfunction in areas such as learning and memory, but little is known about neural mechanisms that contribute to resilience to this adverse effect.

Methods: An existing database of a learning paradigm in sleep deprived and non-sleep deprived 16 to 18-month old C57BL/6 mice was used to identify fast learners and slow learners based on an R2 value representing the learning curve of each individual mouse.

Results: Results showed that sleep deprived mice had more slow learners compared to fast learners whereas non-sleep-deprived mice showed the opposite. Hippocampal immunohistochemistry and digital imaging analysis showed sleep deprived, fast learners expressed lower levels of monocyte chemoattractant protein-1 and histone deacetylase 2 and higher levels of synaptophysin and brain-derived neurotrophic factor compared to sleep-deprived slow learners.

Conclusions: These observations provide evidence to suggest that sleep-deprived mice that performed well in a cognitive assay show less hippocampal mediated learning impairment and provide the rationale for further investigations into neurobiological resilience to sleep deprivation with increasing age.

Keywords: Sleep deprivation, resiliency, learning impairment, aging, neuropathology, hippocampus

Introduction

Environmental adversity can often lead to stress as a biological reaction. The human body reacts to stress by activating the hypothalamic-pituitary-adrenal (HPA) axis and producing stress hormones such as glucocorticoids and cortisol [1]. Dysregulation and chronic exposure to the stress response can then lead to adverse health outcomes. However, stress does not affect everyone in the same way. Individuals in the population who have enhanced stress resilience mechanisms can adapt successfully to stress without the burden of illnesses. Increased resiliency to stress also contributes to a slower aging process and improved overall health and quality of life. Individuals that are less vulnerable to stress and its adverse health effects are deemed resilient [2].

The study of resilience is more common among younger study populations, but less so when it comes to older adults [3]. The aging process itself is one of the most challenging public health issues faced by developed countries. By 2050, the number of people over 65 years old is expected to reach 1.5 billion, nearly tripling the amount in 2010 [4]. This rapid demographic change will put more stress on various social infrastructure systems in countries around the world. With the growth in aged populations, there has been an expansion in initiatives and interventions to promote successful aging [5]. The topic of resilience is a rapidly growing field of interest in research, especially in older adults and the impact it has on successful aging. Aging well has many components, such as being free from chronic disease and maintaining a high level of physical and mental function [6].

Sleep is an essential component of energy conservation and homeostasis of multiple physiological and behavioral processes [7]. The American Academy of Sleep Medicine and the National Sleep Foundation recommend that adults sleep at least 7 hours per day [8,9]; however, the increasing demand for long work shifts and “around-the-clock”
work has led to a marked reduction in the average sleep duration in developed countries [10, 11]. Sleep deprivation leads to numerous cognitive and behavioral effects, such as increased reaction to stress, emotional changes, and a lack of impulse control [12]. The issue of sleep loss has become so pervasive in society that the Center for Disease Control has recently elevated it to public health epidemic status [13]. However, research has also shown that some individuals are vulnerable while others are resistant to sleep deprivation [14, 15]; notably, this phenotypic stability is maintained across months and years [15]. Approximately a third of healthy adults show high levels of decreased performance when moderately sleep deprived. Another third displays moderate deficits while the final third shows little to no decrease in performance even with severe sleep loss. The underlying reasons for such differential neurobehavioral vulnerability to sleep loss are mostly unknown and unexplained by demographic and other factors [16-18]. To date, there have been studies looking at different genetic approaches to relate biomarkers to sleep deprivation responses, although the specific mechanisms for these relationships are currently not fully understood [14].

Phenotypic characteristics of resilience to sleep deprivation should be determined before investigating genetic differences. In this regard, we have developed a mouse model of short-term sleep deprivation and a spatial navigation task (Box maze) designed to assess learning behavior in a time-effective and relatable manner [19]. Therefore, sleep deprivation as a stressor can be used to exploit the cognitive impairments that arise thus facilitating the study of innate resiliency in individual mice [20-22]. In this study, we used an existing database of sleep-deprived mice and analyzed Box maze data from previous studies. We report that sleep-deprived mice that performed well in a Box maze cognitive learning assay show less hippocampal mediated learning impairment, providing the rationale for further investigations into neurobiological resilience with increasing age.

**Methods**

**Box maze database**

Data were used from a Box maze database that had been deposited for over two years on a shared Google drive representing a number of studies from the senior authors’ laboratory. The data used for this report were from 16 to 18-month-old male C57BL/6 (B6) mice provided by the National Institute on Aging Aged Rodent Colony (Charles River, Inc.). After extensive mining of all available data from the drive, a total of n=40 mice were suitable for the hypothesis being tested. Half of the mice underwent learning assessment after being exposed to a short term sleep deprivation protocol while the remaining half were those kept on a standard sleeping cycle. Housing and maintenance have been previously described [19].

**Grading platform**

The box maze produces data in the form of escape latencies with four trials per mouse. This raw data can then be plotted on a graph through programs such as Excel, Graphpad Prism, etc. The grading platform further defines the escape latencies with a single value known as the coefficient of determination (R²). After plotting the raw data, a logarithmic trend line can be applied to the data points in which the R² value and slope are observed for each mouse. If the slope is positive, the mouse is graded as a slow learner whereas mice with a negative slope tend to be fast learners. The logarithmic trend line was chosen as the most appropriate to describe the data points because a learning paradigm should be one in which the data quickly level off [23]. The R² values range from 0 to 1 with 1 being close to the logarithmic trend line.

**Immunohistochemistry**

Immunohistochemistry (IHC) was performed on 5 μm thick, paraffin-embedded mouse brain tissue mounted onto slides. Slides were rehydrated with xylene, decreasing concentrations of ethanol, and deionized water. Antigen retrieval was performed by immersing the slides in a hot water bath at 98 degrees C incubated in a 1:10 Citrate Antigen Retrieval solution in autoclaved deionized water and cooled down to room temperature for 20 minutes. Slides were then stained using an avidin-biotin horseradish peroxidase (HRP) kit (anti-rabbit HRP-3,3’-Diaminobenzidine (DAB) Cell & Tissue Staining Kit, R&D Systems Minneapolis, MN) with manufacturer instructions slightly modified for best staining outcome. Slides were applied with a 3% peroxidase blocking reagent for 15 minutes to quench endogenous peroxidase activity which reduces background noise on the final stain result. Slides were washed in Tris-buffered saline, 0.1% Tween® 20 Detergent (TBST) solution for 5 minutes. To reduce non-specific hydrophobic interactions between the primary antibody and the tissue, a serum blocking reagent was placed onto each section for 15 minutes. After draining the serum off of the slides, avidin blocking reagent was placed onto them for 15 minutes followed by a rinse with TBST for 5 minutes. To prevent the binding of previously applied avidin, biotin blocking reagent was placed onto the slides for 15 minutes. The primary antibody in TBST at the following concentrations: histone deacetylase 2 (HDAC2)1/500 (ab7029, Abcam, Cambridge UK), brain-derived neurotrophic factor (BDNF) 1/500 (ab108319, Abcam, Cambridge UK), monocyte chemoattractant protein-1 (MCP-1) 1/200 (ab25124, Abcam, Cambridge UK), Synaptophysin 1/500 (ab32127, Abcam, Cambridge UK), brain-derived neurotrophic factor 1/500 (ab108319, Abcam, Cambridge UK), brain-derived neurotrophic factor 1/500 (ab108319, Abcam, Cambridge UK) was applied overnight in a humidified chamber. Slides were rinsed 3 times in TBST for 5 minutes each, then incubated with a biotinylated secondary antibody for 30 minutes and rinsed 3 times in TBST for 5 minutes each. Slides were then incubated in High Sensitivity Streptavidin (HSS)-HRP for thirty minutes and rinsed in TBST 3 times for 2 minutes each. DAB Chromogen was applied to slides and incubated in the solution for 5 minutes each before rinsing with
deionized water for 5 minutes. Slides were dehydrated in an increasing concentration of ethanol and xylene then mounted with a coverslip.

**Imaging and QuPath analysis**

IHC slides were photographed under a Nikon Eclipse E400 microscope with a Nikon D7100 camera through a microscope camera adaptor. All photos were taken under a magnification of 4x so that the entire hippocampus could be captured within each photo field. Photos were then uploaded onto a Google drive where slides were separated into project files organized by staining group. QuPath version v0.2.0-m11 was downloaded from Github (https://QuPath.github.io/) [24]. When downloading, QuPath allows the user to determine how much random access memory (RAM) it will take up; so we determined 6 GB would be sufficient for the analysis of the project. A new project was created on QuPath per each staining group examined. This step was done to avoid having to redo the steps of the project workflow for each image. First, the image type was set to hematoxylin and 3,3’-Diaminobenzidine (H-DAB) in order for QuPath to recognize the images as a DAB Chromogen stain. The red, green, and blue (RGB) values for DAB were then calibrated to better represent the project by selecting a region of interest (ROI) that is representative of positive stains along with the hippocampus. Smaller ROIs were used that selected only the smaller areas of positive staining to reduce any potential background noise. These ROIs were then averaged to come up with a new RGB DAB value for the project code. The hippocampus was then annotated with the polygon wand to only measure staining at the desired region of the tissue. However, if there were any folds or staining irregularities through a manual check, the slide was omitted from the analysis. To quantify the staining, superpixels were created to analyze the hippocampus [25]. Within the annotated hippocampus, QuPath groups similar pixels into a cluster called a superpixel based on the RGB values set for DAB. The pixel-based analysis was chosen as the desired method of quantification because this study is looking at multiple stains, and pixel analysis allows us to follow an almost identical protocol between each group [25]. Superpixel size was set to 25 µm$^2$ in order to balance capturing positively stained sections at a high resolution and processing speed. QuPath then applies a DAB intensity to each of the superpixels previously set by the initial DAB RGB calibrating. Not only does it process as positive or negative, but it also separates the staining at three levels of thresholds: 0.2, 0.4, and 0.6 or a positive at all three levels. This, therefore, allowed the capture of staining intensity across all positively stained cells in the annotated region. In order to visualize the DAB staining thresholds, a “heat map” was generated for each image [25]. QuPath allows users to apply a gradient of color according to the quantifications of DAB staining generated previously. The heat map allows a qualitative complement of the previous analyses and serves useful in identifying certain regions that have higher levels of staining. QuPath generates the heat map by assigning a color to each superpixel which indicates the different levels of DAB staining based on the previously described thresholds. However, the upper and lower bounds of the color spectrum must be set by the user and made equal for each new project. This way, the results within each project can be compared to one another relative to staining distribution and intensity across the hippocampus. It should be noted, however, that the color is not indicative of positivity, but rather captures the differences in intensity of the stain.

**Statistics**

Welch’s t-tests, Pearson Correlations, and graph creation were performed using Prism statistical software (Graphpad Software, La Jolla, CA, USA). The p-value was set for a statistical significance of $p<0.05$. All the data were presented as mean ± standard error of mean (SEM).

**Results**

The box maze grading platform separated slow learners from fast learners

Upon analyzing the 20 sleep deprived and 20 control mice that were selected from the database of Box maze results, we found $n=6$ fast learners in the sleep deprived group and $n=15$ fast learners in the control group (Figure 1). Of a total of $n=40$ middle-aged, C57BL/6 mice, $n=20$ were exposed to short-term sleep deprivation whereas $n=20$ were a control group (non-sleep deprived). All $n=40$ were analyzed through the grading platform to classify each mouse as either a fast learner or a slow learner based on the $R^2$ value. In the $n=20$ sleep deprived...
mice, n=6 were classified as sleep deprived, fast learners whereas n=14 were sleep deprived, slow learners. In the n=20 control mice, n=15 were classified as control fast learners whereas n=5 were control, slow learners. Plotting the raw escape latencies for each mouse produced a graph that allowed the further distinction of fast and slow learners through utilizing the coefficient of determination. Fast learners had a sharper and more consistent decrease in latency time over the four trials when compared to slow learners, which can be better modeled with a logarithmic trendline. Thus, fast learners had R² values that were close to 1 while slow learners had R² values close to 0. However, there were also mice with R² values that ranged somewhere in the middle. After finding the average R² value across n=80 control mice, we set the fast learner threshold value at greater than 0.74 (Figure 2).

The results from grading each mouse showed that among the sleep deprived group, more mice had R² values below 0.74 and were designated as slow learners. This meant that fewer mice were able to perform well on the learning assessment, presumably due to the synaptic impairment known to occur with the loss of sleep [26]. As hypothesized, there was a smaller group of fast learners with R² values above 0.74 within the sleep deprived group that were able to maintain higher levels of cognitive functioning despite being exposed to sleep deprivation. The R² values obtained from the grading platform were useful for comparing Box maze data of fast and slow learners with other variables such as sleep deprivation or staining intensity. Because raw Box maze data that is plotted has many factors to account for, such as escape latency and the 4 trials per mouse, it is difficult to compare to other factors. Therefore, the grading platform provides additional value in that it is able to produce a single number to describe the learning curve across the four trials per mouse (Figure 3).

Hippocampal staining thresholds in sleep-deprived fast-learning mice were similar to that of nonsleep-deprived (control) fast-learning mice.

Slides that were stained through immunohistochemistry and imaged were processed through the whole-image analyzing software program, QuPath [24]. The results, expressed in superpixels, showed that HDAC2, MCP-1, Synaptophysin, and BDNF levels in sleep-deprived, fast learning mice were similar to that of the control, fast learning mice. HDAC2 is a histone deacetylase complex which indicates that some level of epigenetic alteration is occurring. HDAC2 functions to repress transcriptional activity and therefore decreases the expression of DNA products in the brain [27]. There are many types of HDAC, but HDAC2 was of interest to our results because it negatively regulates memory formation and synaptic plasticity [28]. Sleep-deprived, fast learning mice had significantly reduced levels of HDAC2 when compared to that of slow learning mice (Figure 4B and C). MCP-1 levels were also significantly reduced in the sleep-deprived fast-learning group which suggests that there was less neuroinflammation occurring in the hippocampus of mice that were graded as fast in their learning when exposed to short-term sleep loss [29]. Synaptophysin and BDNF were of interest to observe synaptic function and plasticity, re-
Figure 4. Positive Superpixel percentage in the hippocampus when stained with MCP-1, HDAC2, synaptophysin, and BDNF. (A) control mice, (B) sleep deprived mice, (C) fast learners, (D) slow learners. *Significant by Welch’s T-Test of control to sleep deprived groups in light of fast and slow learners (A, B) then fast to slow learners in regards to control and sleep deprived mice (C, D).

Figure 5. Sagittal hippocampal sections were imaged at 40x and analyzed using QuPath. Visual representation of positive superpixels using a heat map. Rows represent staining groups while columns show graded groups for control or sleep-deprived mice. Staining intensity is based on a positive superpixel signal with dark blue suggesting lower levels, green being moderate, and red as high. The rows with HDAC2 and MCP-1 show that fast learners had higher amounts of blue signal whereas slow learners showed redder. The rows with Synaptophysin and BDNF showed that fast learners had higher amounts of red signal whereas slow learners showed more blue and green. Adapted in part from Lee et al. [32].
respectively [30, 31]. Because levels of positive superpixels among the sleep-deprived fast learners and control fast learners were similar, these results showed that there are characteristic levels of protein expression in fast learning mice independent of sleep deprivation. The independent t-tests performed also showed that the control group (Figure 4A) is relevant because the statistics are similar to the experimental group. We expected that a portion of the control group should also be slow learners to show the efficacy of the box maze and grading platform. It also shows that the resiliency of the sleep-deprived fast learners was significantly different from the slow learners. Overall, the control group in our study showed similarity to the experimental group in light of the sleep deprivation that was given to the experimental group. The heat map was able to visualize these results in a qualitative way to show that there were higher levels of super positive pixels as indicated by a color gradient (Figure 5) of red being high and blue being low. The heat map also provided additional insight into the distribution of staining at different areas of the hippocampus. Overall, the heat map served as a qualitative understanding of all four strains that were tested through immunohistochemistry. It complemented the superpixel positivity with a quick visual understanding of what was being expressed by the superpixel data.

Learning ability in sleep-deprived fast learners showed a correlation suggestive of a decrease in hippocampal mediated molecular neuropathology.

The superpixel positivity generated from QuPath was graphed against the $R^2$ values obtained from the box maze grading to determine the relationship between escape latency and hippocampal protein expression. Using the Pearson Correlation coefficient and p-value, each staining group was compared to the total percentage of positive superpixels. MCP-1 had a negative Pearson Coefficient of -0.90 and a p value of <0.0001 (Figure 6A) which indicates that there is a strong negative correlation with superpixel positivity as the $R^2$ value increases. This is suggestive of lower inflammation levels in mice that were faster in learning the maze. HDAC2 also had a negative Pearson Coefficient value of -0.84 and a p value of <0.0001 (Figure 6B) which indicates a strong negative correlation of superpixel positivity with decreasing $R^2$ value. Since HDAC2 is a deacetylase complex, this association is suggestive of higher levels of transcriptional repression in DNA products that ultimately impaired learning in mice. Synaptic markers BDNF and Synaptophysin have opposite results in that both showed a strong statistically significant positive correlation in superpixel positivity and $R^2$ value (Figure 6C and D). BDNF has a Pearson coefficient of 0.91 and a p value of <0.0001 which shows that synaptic plasticity according to positive superpixel was increased in fast learning mice. Synaptophysin also showed similar results, as it had a Pearson coefficient of 0.97 and was statistically significant with a p-value of <0.0001. This is indicative of increased synaptic plasticity according to positive superpixel values in mice that are faster learners. The correlations for all listed stains were statistically significant as indicated by their p-values. Direct comparison of individual escape latency values with the individual QuPath superpixel positivity percentage shows that the box maze grading platform and the immunohistochemistry data are consistent with each other. This comparison reveals that fast learners in the maze generally exhibit lower levels of inflammation and histone deacetylation, while synaptic plasticity and function are increased. These results, therefore, offer a statistically significant and novel way to measure levels of neural molecules in mice after being run through the box maze. These escape latencies can then easily be compared with QuPath positive superpixels as a quick way of investigating molecular pathways of learning impairment in the hippocampus.

Discussion

The grading platform for learning in mice is an effective way to separate slow from fast learners in the spatial navigation Box maze. With the $R^2$ threshold set at 0.74 through averaging the escape latencies of n=80 control mice, results of the grading platform produce a single value that can easily be compared to numbers of other variables of interest. Ultimately, the grading gives an $R^2$ value that captures the totality of the Box maze results (escape latency for four trials) for each individual mouse. This platform is also effective in separating fast and slow
learning mice in both short term sleep deprived and control populations that were run through the Box maze. Through this analysis, we found that n=6 mice out of n=20 that were sleep deprived were graded as fast learners in the platform. Control mice, as expected, had results that showed a higher proportion of fast learners n=15 to slow learners n=5.

The results from the grading platform indicate that the separation process yields expected results that can then be used for further investigation of different processes. Through utilizing immunohistochemistry, the results have shown that the grading platform in combination with the short-term sleep deprivation Box maze protocol produces results that are suggestive of a hippocampal learning impairment in slow learning mice. These results can be quickly visualized through a QuPath heat map that shows a difference in superpixel positivity among the different staining groups generated through the grading platform. Although the exact neural mechanisms of sleep deprivation were not investigated in this project, the heat map of the immunohistochemistry stains provides evidence of positive staining in regard to location and intensity based on the novel idea of super pixels through QuPath, and provides a glimpse of the molecular pathway involved. QuPath is a new software so there is still much to be explored and refined with analyzing protocols. Currently, a PubMed search only brings up a total of 22 papers in total that have used this program to analyze whole image slides. Additionally, because QuPath is only an analyzing software, it is critical that immunohistochemistry protocols be performed in a uniform and consistent manner[32]. This would require careful planning of tissue harvesting, sectioning, and imaging that would allow for the best results. Further molecular analysis should be performed to confirm the QuPath data so that conclusions can be drawn on the molecular pathway associated with hippocampal learning impairment observed in sleep deprived mice. QuPath was able to show a significant correlation between positive super pixels and box maze learning times through the R² value. These observations are therefore a valuable insight into utilizing the Box maze learning classification of slow or fast learners in future studies looking into resilience to environmental stressors of sleep deprivation. QuPath can also be used as a novel application to study correlations between positive super pixels in immunohistochemistry slides from a variety of different protocols and tissues.

We looked at four antibodies in this study: MCP-1, HDAC2, Synaptophysin, and BDNF. Through QuPath analysis, we found that levels of MCP-1 and HDAC increase, while levels of Synaptophysin and BDNF decrease as learning latency increases. Future studies to determine if specific genes were over or underregulated to cause these differences could help elucidate the genetic pathways involved in resilience. Furthermore, it would be interesting to see if there were some genetic differences between slow and fast learning mice that predisposed them to learning at their respective speeds. This could shed light on genetic predisposition to resilience and point to targets that could be studied in humans.

The model of short-term sleep deprivation in middle age is translationally relevant because older adults are observed to have reduced sleep according to electroencephalography (EEG) data in comparison to younger adults[33]. Mice 16-18 months old represent the general human population in their late 50’s and early 60’s[34]. This is an important age group to study resilience, as cognitive decline is often seen shortly after midlife and occurs noticeably at ages 70 or higher[35]. Our study found significant differences in four protein levels between fast and slow learning mice. The data generated in this study suggest that mice can be used as models for sleep deprivation and resilience and offer valuable insight on the importance of intervention to attenuate the impacts of cognitive decline that is often observed in the older population.

Declarations

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