Near infra-red light treatment of Alzheimer’s disease

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Alzheimer’s disease (AD) is a chronic neurodegenerative disease. The symptoms include memory and spatial learning difficulties, language disorders, and loss of motivation, which get worse over time, eventually ending in death. No effective treatments are available for AD, currently. Current treatments only attenuate symptoms temporarily and are associated with severe side effects. Near infra-red (NIR) light has been studied for a long time. We investigated the effect of NIR on AD using a transgenic mouse model, which was obtained by co-injecting two vectors carrying AD mutations in amyloid precursor protein (APP) and presenilin-1 (PSEN1) into C57BL/6J mice. The irradiation equipment consisted of an accommodating box and an LED array. The wavelength of NIR light emitted from LED was between 1040 nm and 1090 nm. The power density delivered at the level of the mice was approximately 15 mW/cm². Firstly, we treated the mice with NIR for 40 days. Then, the irradiation was suspended for 28 days. Finally, another 15 days treatment was brought to mice. We conducted Morris water maze and immuno-fluorescence analysis to evaluate the effects of treatment. Immuno-fluorescence analysis was based on measuring the quantity of plaques in mouse brain slices. Our results show that NIR light improves memory and spatial learning ability and reduces plaques moderately. NIR light represents a potential treatment for AD.

Keywords: Alzheimer’s disease; near infra-red light; transgenic mouse model; Morris water maze; immuno-fluorescence analysis.

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1. Introduction

Alzheimer’s disease (AD) tends to occur in people aged above 65 years. It is a neurodegenerative disease and difficult to cure. Current therapeutic methods, mainly including medications, have limited effects in attenuating symptoms. No treatment can reverse the progression of disease. The pathogenesis of AD is still poorly understood. It is believed that amyloid-β (Aβ) deposition leads to the incidence and deterioration of AD. Aβ is derived from the regulated intracellular proteolysis of amyloid precursor protein (APP), which extensively exists in tissues. APP is hydrolyzed by α-, β-, and γ-secretase. Hydrolysis of APP by β-, and γ-secretase leads to Aβ formation. Aβ peptides contain 39–43 amino acids each, and Aβ1–40 and Aβ1–42 are the major peptides associated with AD plaque formation via aggregation. Aβ and senile plaques have long been considered neurotoxic and associated with the progression of AD. Plaque reduction is the therapeutic goal currently.

Light has been used for medical applications for a very long time. Near infra-red (NIR) light, a section of electromagnetic spectrum, has good performance in detection and treatment. The determination of NIR spectroscopy is useful for quantitative analysis of glycated hemoglobin and biomedical imaging in living tissues. NIR is therapeutic, due in part to its deep tissue penetration and low light scattering. NIR is typically applied with its thermal effects or biological effects in treatment. The heat converted from NIR helps the release of drug in microcapsule system. Biological effects of light include wound healing, pain reduction, and alleviation of oral mucositis. Effects such as cytoprotection, cellular proliferation, growth factor release have also been reported. Studies have indicated the effects of NIR on nervous system, like AD. The biological effects of NIR are specific to wavelength and not based on thermal effects. Those studies showed NIR mitigated AD neuropathology and behavioral deficits. For example, the CD1 mouse model treated with 1072 nm low infra-red light shows improvements in emotional responses and memory performance. AD mice which had undergone five month treatment with 1072 nm NIR showed a significant reduction of Aβ1–42 in the cerebral cortex and up-regulation of stress response proteins in the brain, which were known to reduce protein aggregation and neuronal apoptosis.

The 808 nm wavelength significantly reduced amyloid plaques and improved behavioral performance in AD mouse model. Purushothuman et al. applied 670 nm NIR to two kinds of transgenic mouse models, finding out that NIR was associated with a reduction in hyperphosphorylated tau, neurofibrillary tangles, oxidative stress markers and Aβ plaques in the neocortex and hippocampus.

We used a transgenic AD mouse model to evaluate the effect of NIR. The mouse model was co-injected with two mutant gene vectors involving APP and PSEN1. APP and PSEN1 mutations increase Aβ formation. Our transgenic model exhibits plaques and memory deficits at an early age. In this present study, we treated the AD mouse model using NIR light and analyzed the effects through Morris water maze (MWM) and immunofluorescence analysis (IFA). We chose different irradiation parameters from previous reports and designed a suspension between two periods of irradiation to assess the sustainability of the effect of NIR.

2. Materials and Methods

2.1. Animals

All our mice were obtained from the Institute of Laboratory Animal Science, Chinese Academy of Medical Sciences. The transgenic mouse model used in this study was produced by co-injecting the APPsw and PS1ΔE9 vectors. The APPsw vector expresses a mouse–human hybrid transgene containing the extracellular and intracellular regions of the mouse sequence as well as a human sequence within the Aβ domain with Swedish mutations (K594N/M595L). The PS1ΔE9 vector expresses the human presenilin-1 deleted exon-9. The background strain of the mice was C57BL/6J. The transgenic model showed memory and spatial learning disability as well as detectable plaques at an early age. Senile plaques were detected as early as 4.5 months. And behavioral differences between the model and wild-type mice appeared in three months. All the animals were female and divided into three different groups ad (n = 10), ad + ir (n = 10) and wt (n = 12). Group ad was transgenic mice and did not received irradiation. Group ad + ir was transgenic mice and received irradiation. Group wt was normal mice and did not received irradiation. All the mice were raised under 12 h light/12 h dark cycle, and exposed to light from 7 am to 7 pm.
The room temperature was maintained constant at 23 ± 2°C. Mice were provided free access to water and food, and their cages were cleaned twice a week. Animal care and experimental protocols were in accordance with guidelines issued by the Shanghai Medical Experimental Animal Care Commission. At the start of the MWM task, we found four and two dead mice in the ad + ir and ad groups, respectively. Mice in the ad + ir group died because of disrupted air supply, while mice in other groups were not affected. Two ad mice were attacked by others in the same cage. All the six mice died, but not because of IR irradiation. This study was approved by the Ethical Committee of Animal Experiments of Med-X Research Institutes, Shanghai Jiao Tong University.

2.2. Methods and treatment

Our patented apparatus (ZL 2015 2 0692323.1) consists of a box containing mice and an LED array, which emits NIR light. The LED array also acts as the lid. The side windows and vertical openings provide air, food and water. We irradiated the ad + ir group of mice six min per day for 40 days, at the age of six months, with a power density of approximately 15 mW/cm² and a wavelength between 1040 nm and 1090 nm. We chose 6 min according to previous studies. Then, we stopped the treatment and tested the memory and spatial learning ability of all the mice using a Morris water maze. The time taken by the mouse to locate the platform successfully within 60 s was recorded as the latency time. Otherwise, the latency time was counted as 60 s. We consider the trial complete when the mouse located the platform or after a lapse of 60 s. The mouse was relocated at the platform for 15 s to acclimatize with the surroundings. Each mouse was tested over four trials consecutively, daily. The software we used to record data was ANY-maze 4.99 (Stoelting Co.).

2.4. Immunofluorescence analysis

All the mice were euthanized via CO₂ narcosis by rapidly dissecting the brain. The brains were

<table>
<thead>
<tr>
<th>Group</th>
<th>N before treatment</th>
<th>First irradiation</th>
<th>First MWM</th>
<th>Irradiation suspension</th>
<th>Second MWM</th>
<th>Second irradiation</th>
<th>N after treatment</th>
<th>IFA</th>
</tr>
</thead>
<tbody>
<tr>
<td>ad + ir</td>
<td>10</td>
<td>1–40</td>
<td>41–47</td>
<td>41–68</td>
<td>69–73</td>
<td>74–88</td>
<td>8</td>
<td>Yes</td>
</tr>
<tr>
<td>ad</td>
<td>10</td>
<td>No</td>
<td>41–47</td>
<td>No</td>
<td>69–73</td>
<td>No</td>
<td>6</td>
<td>Yes</td>
</tr>
<tr>
<td>wt</td>
<td>12</td>
<td>No</td>
<td>41–47</td>
<td>No</td>
<td>69–73</td>
<td>No</td>
<td>12</td>
<td>Yes</td>
</tr>
</tbody>
</table>
immediately frozen in iso-pentane and stored at −80°C. Brains were cut into coronal sections of 20–μm thickness at −20°C. Endogenous peroxidase activity was inhibited by incubating the sections in 0.3% peroxide in methanol for 20 min at room temperature. The samples were subjected to three 5-min washes with phosphate buffer saline (PBS). The sections were fixed with 4% paraformaldehyde (PFA) in PBS (pH7.4) for 15 min and washed for 5 min three times, again. Subsequently, the sections were blocked with 10% normal donkey serum in 0.1% Triton X-100 for 30 min and incubated with the primary antibody (Abcam, ab2539, 1:100) in 1% normal donkey serum overnight at 4°C. The primary antibody was sensitive to synthetic peptides corresponding to human beta amyloid aa 1–14 and reacted with all types of Aβ. Before incubation with a second antibody, the samples were washed for 5 min three times. The anti-rabbit second antibody (Jackson, anti-rabbit 488 nm, 1:400) was used in 1% normal donkey serum for 1 h at room temperature followed by a 5-min wash three times. The slices were mounted using a mounting medium with DAPI (Vector laboratories, H-1200), commonly used for nuclear staining. Finally, the sections were imaged by the Leica fluorescence microscope at an excitation wavelength of about 360 nm to locate the nuclei in the hippocampus. Then, we switched the wavelength to 488 nm to count the plaques.

We analyzed the hippocampus and cerebral cortex of mouse brain for immunofluorescence because Aβ was mainly deposited in these areas.26 The plaques within the regions were divided into two categories depending on the counting sizes: small (< 20 μm) and large (≥ 20 μm). Data were derived from three pictures from each mouse held in a fixed position of hippocampus to compare different mice. Plaques were counted manually and repeated three times. The counting was blinded to experimental paradigms. The number of plaques indicate the degree of AD severity. Therefore, we counted the number of plaques in the three groups to assess the effects of NIR irradiation therapy.

2.5. Statistical analysis

The data of each mouse in MWM were averaged and expressed as mean ± SEM. We analyzed the data with SPSS to compare the means of different groups using two-way ANOVA, with group representing the between-subject factor, and time as the within-subject factor. LSD post-hoc analysis was used to evaluate the differences between ad + ir and the other two groups. In IFA, we analyzed the data through two-way ANOVA with Graphpad followed by Bonferroni post-hoc analysis.

3. Results

The results of the first round MWM are presented in Fig. 1. Figure 1(a) shows the column graph of the first round MWM data. Overall, the daily duration of the three groups appeared in the same sequence, from large to small. The ad group lasted longer than the ad + ir group. And the latency time of ad + ir group was longer than that in wt. Significant differences were found between ad and ad + ir groups on days 2 and 3 while significant differences between ad + ir and wt groups were observed on days 2, 3 and 4. The time that the mouse spent in the MWM

![Fig. 1. The results of the first round MWM task. (a) The column graph of first round MWM data. (b) The line chart of first round Morris water data. All data are expressed as means ± SEM. We compared the means of ad (n = 8) and wt (n = 12) with that of ad + ir (n = 6) using two-way ANOVA followed by LSD post-hoc (*p < 0.05, **p < 0.01, ***p < 0.001).](image-url)
task indicated the level of its memory and spatial learning ability. More capable the mouse was, the smaller the value of latency time was. Therefore, the constant sequence showed that the ad + ir mice performed better than the ad mice, which was worse than mice in wt group. The average time of group ad + ir was 44.8 s, which was 7.4% less than the average time of group ad and 18.5% more than the average time of group wt. The two-way ANOVA showed p value between ad and ad + ir of group factor was 0.0049. Figure 1(b) shows the line chart of the first round MWM. The line connects the means of daily data and reveals the variation trend in latency time. The height of the line represents the overall performance of the group. Two lines were separated from each other, clearly showing the differences in their means. All the results indicated that mice in the ad + ir group performed better than those in ad after phototherapy, suggesting that the NIR light might be effective for AD treatment.

Figure 2 shows the results obtained from the second round of MWM after 28 days of treatment suspension. As shown in Fig. 2(a), the column graph of the second round MWM revealed no significant difference between ad and ad + ir groups. Figure 2(b) presents the line chart of the second round MWM. The ad and ad + ir groups were almost similar and intertwined with each other. The second-round MWM data suggested that after 28 days of light therapy suspension, the ad + ir group performance was almost same to that of ad, which was not irradiated. Therefore, these results indicated that the effects of IR treatment disappeared over time.

After the behavioral experiments, we used biochemical techniques to analyze the secretion of Aβ by labeling the plaques and counting their numbers in mouse brains. Immunofluorescence analysis was conducted according to standard procedures. The primary antibody used was sensitive to the synthetic peptide corresponding to human beta-amyloid amino acids 1–14 conjugated to Keyhole Limpet Haemocyanin (KLH). An interesting study showed that both newly formed and existing plaques grew at a similar rate of 0.3 μm (radius) per week.27 We assumed that IR treatment decreased amyloid plaques by interfering with the formation of new plaques. Therefore, we divided the plaques into two categories depending on their size: small (<20 μm) and large (≥20 μm).

The results of IFA are shown in Fig. 3. Figures 3(a) and 3(b) illustrate immunofluorescence results. Figure 2(c) shows the column graph of ad (n = 4) and ad + ir (n = 3) data. No plaques were found in wt mouse brain. As shown in Fig. 3, mice in the ad + ir group obviously showed fewer plaques than those in the ad group, with an average number of 15.33 and 35.5 in ad + ir and ad groups, respectively. Due to small sample size, the variance was not adequate to generate significant differences between the two groups. The mean number of plaques measuring <20 μm was 9 and 26 in ad + ir and ad groups, respectively. And the average number of plaques measuring ≥20 μm was 11 and 6.3 in ad and ad + ir groups, respectively. NIR treatment modestly reduced the small-size plaques, which was consistent with a previous study.19 There was significant difference (p < 0.05) between the two groups for plaques measuring <20 μm. And for plaques measuring ≥20 μm, p value was bigger than 0.05. The small sample size and short duration of irradiation might contribute to large p value. More mice need to be tested in future studies.
4. Discussion and Conclusion

Michalikova et al. administered 1072 nm infra-red light to CD1 mice to test their emotional response and memory performance. Ten days of exposure improved the working memory of mice.16 Another study reported by Grillo also employed 1072 nm to treat a transgenic mouse model over five months.17 In contrast to this chronic and low-power treatment protocol with 5 mW/cm², we adopt a daily therapy using a higher power density with about 15 mW/cm². Purushothuman et al. studied two mouse models with NIR treatment, and found out that NIR treatment could reduce hyperphosphorylated tau, neurofibrillary tangles, oxidative stress markers and plaques.19 They then extended the irradiated area to cerebellum. After one month of treatment, NIR mitigated AD-related pathologies.28 The mechanisms associated with NIR effect on AD mouse are typically mediated via ATP generation to increase and improve mitochondrial function.29 Taboada et al. used 808 nm in AβPP transgenic mice, which increased the ATP levels as well as percent oxygen consumption.18 Previous reports suggest that early stage plaques could be more toxic than those at later stages.30 The early damage essentially involved cellular processes resulting in cell death. NIR treatment diminished cell death associated with amyloid plaque formation in transgenic mice, resulting in improved spatial learning and memory.

Though the NIR we used penetrated tissues deeply compared to other wavelengths,31 the unshaved fur of mice is still a major limitation because black fur blocks a major portion of the light. Considering the fur and skull of humans are thicker than those of mice, the power density could be higher. Shaving hair may help improve the penetration of NIR and the therapeutic effect. In addition, our conclusions were mainly based on Aβ deposits. Additional histological endpoints are needed considering the limitations associated with a
single standard. Those histological endpoints include phosphorylated tau protein, heat-shock protein, inflammatory markers and so on. In our subsequent study, another group of ad + ir which receive 40 day irradiation only could be included to check the development of plaques after the 40 day period and also to eliminate the confounding effects of the two separate NIR treatments.

Our study applied different irradiation parameters and designed a suspension between two periods of irradiation. Results showed that the parameters we chose worked well and the NIR effect was not permanent. Both provided important insights into clinical application of NIR on treating AD.

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