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Cellular Source-Specific Effects of Apolipoprotein (Apo) E4 on Dendrite Arborization and Dendritic Spine Development

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Abstract

Apolipoprotein (apo) E4 is the leading genetic risk factor for Alzheimer’s disease (AD), and it has a gene dose-dependent effect on the risk and age of onset of AD. Although apoE4 is primarily produced by astrocytes in the brain, neurons can also produce apoE4 under stress conditions. ApoE4 is known to inhibit neurite outgrowth and spine development in vitro and in vivo, but the potential influence of apoE4’s cellular source on dendritic arborization and spine development has not yet been investigated. In this study, we report impairments in dendritic arborization and a loss of spines, especially thin (learning) and mushroom (memory) spines, in the hippocampus and entorhinal cortex of 19–21-month-old female neuron-specific-enolase (NSE)-apoE4 and apoE4-knockin (KI) mice compared to their respective apoE3-expressing counterparts. In general, NSE-apoE4 mice had more severe and widespread deficits in dendritic arborization as well as spine density and morphology than apoE4-KI mice. The loss of dendritic spines, especially mushroom spines, occurred in NSE-apoE4 mice as early as 7–8 months of age. In contrast, glial fibrillary acidic protein (GFAP)-apoE4 mice, which express apoE4 solely in astrocytes, did not have impairments in their dendrite arborization or spine density and morphology compared to GFAP-apoE3 mice at both ages. These results indicate that the effects of apoE4 on dendrite arborization, spine density, and spine morphology depend critically on its cellular source, with neuronal apoE4 having more detrimental effects than astrocytic apoE4.

Introduction

Apolipoprotein (apo) E is a 299-amino-acid glycoprotein that is highly expressed in the mammalian brain and occurs as three isoforms in humans- apoE2, apoE3, and apoE4. Although these isoforms differ from each other by only one or two amino acid substitutions at positions 112 and 158, they have profound functional differences in relation to neurological disorders [1,2,3,4]. ApoE4 is the major genetic risk factor for Alzheimer’s disease (AD) [5,6], with a gene dose-dependent effect on the risk and age of disease onset. Individuals with one copy of the apoE4 gene have a 45% chance of developing AD by age 85, which increases to 50–90% for individuals with two copies [7,8].

ApoE is primarily synthesized by astrocytes in the brain, but it can also be made by neurons under stress conditions [3,9,10,11]. The cellular source of apoE4 influences its pathological effects in vitro and in vivo. For instance, neuronal – but not astrocytic – apoE4 undergoes proteolytic cleavage, resulting in C-terminally truncated fragments that cause tau phosphorylation, mitochondrial impairments, and neurodegeneration [3,12,13,14]. Neuronal apoE4 also increases susceptibility to excitotoxin-induced cell death compared to neuronal apoE3, whereas astrocytic apoE3 and apoE4 are equally excitoprotective [15]. Neuron-specific-enolase (NSE)-apoE4 mice, which express human apoE4 solely in neurons, exhibit spatial learning and memory deficits compared to NSE-apoE3 mice [16]. ApoE4-knockin (KI) mice, which express apoE4 under the control of an endogenous apoE promoter, are also impaired in spatial learning and memory, albeit at an older age [17]. In contrast, mice whose apoE4 expression is limited to astrocytes do not show spatial learning and memory deficits, although they do show impairments in working memory [18].

Dendritic pathology and the loss of dendritic spines and synapses are prominent features in early-stage AD [19,20]. ApoE4-associated abnormalities in dendrite arborization and spine development were reported in both primary neuronal cultures and in transgenic mice [21,22,23,24,25,26], which has been suggested to be linked to apoE4- and its fragment-induced mitochondrial impairments and an inherited variable ploy-T repeat genotype in the mitochondrial out-membrane protein TOMM40 [3,24,27,28]. However, whether these effects are influenced by the cellular source of apoE4 has not yet been investigated. Here, we report impaired dendrite arborization and
Figure 1. Cellular source of apoE influences dendrite number and length in the CA1 region of the hippocampus at 19–21 months of age. (A) A representative Golgi-stained image of a CA1 pyramidal neuron. (B) Quantification of the total number of CA1 dendrites, combining AO+BS dendrites. (C) Number of AO and BS dendrites in the CA1. (D) Quantification of the total length of CA1 dendrites, combining AO+BS dendrites. (E) Length of AO and BS dendrites in the CA1. Values are mean ± SEM, n = 13–15 neurons from 3–5 mice for each group. *p<0.05.

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decreased total, thin, and mushroom spine densities in female apoE4-KI and NSE-apoE4 mice compared to their apoE3 counterparts in both the hippocampus and the entorhinal cortex at 19–21 months of age. In contrast, the dendritic arborization and spine density of glial fibrillary acidic protein (GFAP)-apoE4 mice— which express apoE4 solely in astrocytes— were not impaired compared to GFAP-apoE3 mice. These results indicate that the cellular source of apoE4 profoundly influences its effects on neuronal structure.

Materials and Methods

Transgenic Mice

We analyzed transgenic mice expressing human apoE3 or apoE4 in CNS neurons under control of a neuron-specific enolase promoter on a mouse apoE-deficient background [15,16,29]. All mice were backcrossed >20 times with apoE-deficient mice (C57BL/6J-Apoetm-1Unc; The Jackson Laboratory, Bar Harbor, ME). Transgenic mice expressing human apoE3 or apoE4 driven by a glial fibrillary acidic protein promoter (backcrossed >10 times) were generated on a mouse apoE-deficient background as described [13,15]. Human apoE3- or apoE4-KI mice, in which the mouse apoE gene is replaced by the human apoE sequence [30], were from Taconic. All animal procedures were approved by the Gladstone Institutes and the University of California San Francisco Animal Care and Use Committees.

Golgi Staining

Golgi staining was performed on all mouse genotypes at 7–8 and 19–21 months of age (n = 3–5 mice/genotype). For these experiments, we used the FD Rapid Golgi Stain kit (FD NeuroTechnologies, Ellicott City, MD) as described previously [26,31]. Briefly, freshly dissected brains were immersed in equal parts solution A and B for 2 weeks at room temperature, then transferred to solution C for 4 days at 4°C. Brains were sliced at a thickness of 150 μm using a freezing-sliding microtome (Leica SM2000R).

Analyses of Dendrite Arborization and Dendritic Spine Density and Morphology

To analyze dendrite length and number, brightfield images of individual pyramidal neurons were taken with a Leica CTR5000 10X objective. Apical and basal dendrites were traced using the NeuroJ plugin of ImageJ [32], and total dendrite length and number were recorded. For analysis of dendritic spine density and morphology, the apical and basal dendrites of pyramidal neurons from the CA1 region of the hippocampus, entorhinal cortex, and auditory cortex were imaged with a Leica CTR5000 brightfield 63X oil objective. Protrusions were classified into one of four subtypes—stubby, thin, mushroom, or filopodia—based on published criteria [31,33]. Images were coded and analyzed in a blinded manner using ImageJ software [24,31,34].

Statistical Analysis

All values are expressed as mean ± SEM. Statistical analyses were performed with GraphPad Prism software. Differences between the means were assessed by t-test or one-way ANOVA. A p-value of <0.05 was considered to be statistically significant. Statistical values are denoted as follows: *p<0.05, **p<0.01, ***p<0.001.

Results

The Cellular Source of apoE4 Determines its Effects on Dendritic Arborization

We examined dendritic arborization in several apoE4 mouse models and their apoE3-expressing controls; namely: NSE-apoE4 (versus NSE-apoE3), apoE4-KI (versus apoE3-KI), and GFAP-apoE4 (versus GFAP-apoE3). Since some of these apoE4 mice did not develop spatial learning and memory deficits until 16–18 months [17], we first analyzed these mice at 19–21 months of age. Only female mice were used due to their enhanced susceptibility to apoE4-induced learning and memory deficits [16,17,35]. Pyramidal neurons from the CA1 region of the hippocampus, the entorhinal cortex, and the auditory cortex were analyzed using the rapid Golgi-Cox impregnation method. Apical oblique (AO) and basal shaft (BS) dendrites were examined separately.

We first measured the total length and number of CA1 dendrites in NSE-apoE4, apoE4-KI, and GFAP-apoE4 mice and their respective apoE3 counterparts (Fig. 1A). Quantitative analysis revealed that in the CA1, the total number of dendrites in NSE-apoE4 mice was higher than in NSE-apoE4 mice (Fig. 1B). Further analysis revealed a trend toward a greater number of apical dendrites in NSE-apoE3 mice than in NSE-apoE4 mice in the CA1, although this difference did not reach statistical significance (Fig. 1C). NSE-apoE4 mice showed a trend toward decreased total dendrite length in the CA1 (Fig. 1D) compared to NSE-apoE3 mice, and basal NSE-apoE4 dendrites were significantly shorter than basal NSE-apoE3 dendrites (Fig. 1E). One-way ANOVA revealed significantly shorter basal dendritic length in NSE-apoE4 mice than in apoE4-KI and GFAP-apoE4 mice (Fig. 1E). Neither apoE4-KI nor GFAP-apoE4 mice showed significant differences in dendrite number (Fig. 1B) or length (Fig. 1D) compared to their apoE3 controls.

As in the CA1, NSE-apoE4 mice had a greater number of dendrites in the entorhinal cortex than NSE-apoE4 mice (Fig. 2A,B), and this difference reflected changes in basal dendrite numbers (Fig. 2C). A reduction in total dendritic length was also observed in NSE-apoE4 mice (Fig. 2D). In contrast, apoE4-KI and GFAP-apoE4 mice had similar dendritic numbers (Fig. 2B) and lengths (Fig. 2D) compared to their apoE3 controls in the entorhinal cortex. When analyzed separately, apical, but not basal, dendrites showed a trend toward shorter length in NSE-apoE4 mice compared to NSE-apoE3 mice (Fig. 2E). In the auditory cortex (Fig. 3A), GFAP-apoE4 mice showed an increase in total dendritic number (Fig. 3B) and length (Fig. 3D) compared to GFAP-apoE3 mice, which reflected changes in apical, but not basal, dendrite number (Fig. 3C,E). However, neither apoE4-KI nor NSE-apoE4 mice showed differences in dendrite number or length compared to their apoE3 controls in the auditory cortex (Fig. 3B,D).

The Cellular Source of apoE4 Determines its Effects on Dendritic Spine Density

We next examined whether the cellular source of apoE4 might differentially impact dendritic spine density. In the CA1 region, both NSE-apoE4 and apoE4-KI mice had significant reductions in total spine number compared to their apoE3 controls. Surprisingly, GFAP-apoE4 mice had more spines compared to GFAP-apoE3 mice in the CA1 region (Fig. 4A,B). Moreover, these differences exclusively reflected alterations in basal spines, with the apical spine densities showing no significant differences (Fig. 4C). One-way ANOVA revealed significant reductions in total and basal spines in NSE-apoE4 and apoE4-KI mice compared to GFAP-apoE4 mice (Fig. 4B,C).
Figure 2. Cellular source of apoE influences dendrite number and length in the entorhinal cortex at 19–21 months of age. (A) A representative Golgi-stained image of a pyramidal neuron from the entorhinal cortex. (B) Quantification of the total number of entorhinal cortex dendrites, combining AO+BS dendrites. (C) Number of AO and BS dendrites in the entorhinal cortex. (D) Quantification of the total length of entorhinal cortex dendrites, combining AO+BS dendrites. (E) Length of AO and BS dendrites in the entorhinal cortex. Values are mean ± SEM, n = 13–15 neurons from 3–5 mice for each group. *p < 0.05.

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Figure 3. Cellular source of apoE influences dendrite number and length in the auditory cortex at 19–21 months of age. (A) A representative Golgi-stained image of a pyramidal neuron from the auditory cortex. (B) Quantification of the total number of auditory cortex dendrites, combining AO+BS dendrites. (C) Number of AO and BS dendrites in the auditory cortex. (D) Quantification of the total length of auditory cortex dendrites, combining AO+BS dendrites. (E) Number of AO and BS dendrites in the auditory cortex. Values are mean ± SEM, n = 13–15 neurons from 3–5 mice for each group. *p<0.05, **p<0.01.

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In the entorhinal cortex, similar decreases in total spine density were also evident in NSE-apoE4 (versus NSE-apoE3) and apoE4-KI (versus apoE3-KI) mice (Fig. 5A,B). Further classification of apical and basal spines revealed that in apoE4-KI mice, only the apical spine numbers were decreased, whereas NSE-apoE4 mice had fewer spines along both apical and basal dendrites (Fig. 5C). Astrocytic apoE expression did not affect total spine densities in either the apical or basal neuronal regions, as these densities were highly similar between GFAP-apoE4 and GFAP-apoE3 mice (Fig. 5A–C). One-way ANOVA revealed significant reductions in total, apical, and basal spines in NSE-apoE4 mice compared to GFAP-apoE4 mice (Fig. 5B,C).

In the auditory cortex, NSE-apoE4 mice showed a significant decrease in spine density compared to apoE3 controls (Fig. 6A–C). Both NSE-apoE4 and GFAP-apoE4 mice had fewer basal mushroom spines compared to apoE4-KI mice (Fig. 6C), whereas GFAP-apoE4 mice had more mushroom spines than GFAP-apoE3 mice (Fig. 6E), and this difference was reflected in basal but not apical spine density (Fig. 6F). One-way ANOVA revealed significant reductions in total and basal spine densities in NSE-apoE4 mice compared to apoE3-KI mice (Fig. 6B,C).

The Cellular Source of apoE4 Determines its Effects on Dendritic Spine Morphology

Since dendritic spine morphology reflects synaptic function and plasticity [36], we also compared the numbers of spine subtypes in different brain regions of the apoE3 or apoE4 mice. Dendritic protrusions were classified into four categories based on the head size and neck length: stubby, thin, mushroom, and filopodia [31,34]. All data were summarized in Table 1. In the CA1 region of the hippocampus, we observed decreases in the total number of stubby (Fig. 7A) and thin (Fig. 7C) spines in NSE-apoE4 compared to apoE3-KI mice. Selective examination of the basal dendrites revealed a significant drop in stubby spines in apoE4-KI mice as well as NSE-apoE4 mice compared to their apoE3 counterparts (Fig. 7B), whereas thin spines were only reduced in NSE-apoE4 mice compared to NSE-apoE3 mice. ApoE4-KI mice had fewer basal mushroom spines compared to apoE4-KI mice (Fig. 7F), whereas GFAP-apoE4 mice had more mushroom spines than GFAP-apoE3 mice (Fig. 7E), and this difference was reflected in basal but not apical spine density (Fig. 7F). One-way ANOVA revealed significant reductions in total and basal spine densities in apoE4-KI and NSE-apoE4 mice compared to apoE3-KI mice (Fig. 7F).

In the entorhinal cortex, thin spine density was decreased in apoE4-KI compared to apoE3-KI mice (Fig. 8C), which involved the apical, but not basal, spine subset (Fig. 8D). NSE-apoE4 mice...
had fewer total (Fig. 8E) and basal (Fig. 8F) mushroom spines compared to NSE-apoE3 mice in this region. One-way ANOVA revealed significant reductions in total and basal mushroom spines in NSE-apoE4 mice compared to GFAP-apoE4 mice (Fig. 8E,F). The densities of stubby spines (Fig. 8A–B) and filopodia (Fig. 8G–H) did not differ between any pairs in the entorhinal cortex. Finally, in the auditory cortex, NSE-apoE4 mice showed a reduction in total (Fig. 9E) and basal (Fig. 9F) mushroom spine densities compared to NSE-apoE3 mice. One-way ANOVA revealed significant reductions in total and basal mushroom spines in NSE-apoE4 and GFAP-apoE4 mice compared to apoE4-KI mice (Fig. 9E,F). The number of filopodia was higher in GFAP-apoE4 mice compared to GFAP-apoE3 mice (Fig. 9G–H). There were no significant differences in stubby (Fig. 9A–B) or thin (Fig. 9C–D) spine densities between any pairs in the auditory cortex.

The Dendritic Spine Loss Occurs in NSE-apoE4 Mice at Young Ages

We previously reported that NSE-apoE4 mice had learning and memory deficits at 7–8 months of age [13,15,16], whereas apoE4-KI and GFAP-apoE4 mice did not have similar deficits at these ages [17,18]. To explore the potential relationship between apoE4-induced dendritic spine loss and behavioral deficits, we also assessed the dendritic arborization and spine density and morphology in different brain regions of the apoE3 or apoE4 mice at 7–8 months of age. Although apoE4 had no significant effects on dendritic arborization in all three models of mice at these young ages (data not shown), it significantly decreased total spine density in the entorhinal cortex of NSE-apoE4 mice compared to NSE-apoE3 mice (Fig. 10B). ApoE4 also significantly decreased total mushroom spines in the CA1 subregion of the hippocampus, the entorhinal cortex, and the auditory cortex of NSE-apoE4 mice compared to NSE-apoE3 mice (Fig. 10D–F). One-way ANOVA revealed significant reductions in total mushroom spines in NSE-apoE4 mice compared to apoE4-KI and GFAP-apoE4 mice in the CA1, the entorhinal cortex, and the auditory cortex (Fig. 10D–F). ApoE4-KI and GFAP-apoE4 mice had no differences in their spine densities in all three regions of the brain compared to their respective apoE3-expressing counterparts (Fig. 10D–F).

Discussion

In this study, we used mice expressing human apoE to determine whether the effects of apoE on dendritic morphology depend, at least in part, on the cell type in which it is synthesized. Indeed, in female apoE mice at 19–21 months of age, we report
cellular source-dependent effects of apoE4 on dendrite arborization as well as spine density and morphology in vivo. Whereas NSE-apoE4 and apoE4-KI mice show dendritic arborization deficits and/or spine loss in the hippocampus and different regions of the cortex compared to their respective apoE3 counterparts, GFAP-apoE4 mice do not have such impairments. Furthermore, of the mouse models investigated, NSE-apoE4 mice had the most severe and widespread morphological deficits, followed by apoE4-KI mice. The loss of dendritic spines, especially mushroom spines, occurred in NSE-apoE4 mice as early as 7–8 months of age. These data suggest that neuron-derived apoE4 is especially neurotoxic, with profound effects on dendrite arborization, spine density, and spine morphology, while astrocyte-derived apoE4 does not have these detrimental effects.

The Effect of apoE4’s Cellular Source on Dendrite Development

Previous studies have shown that apoE3 promotes neurite outgrowth and dendritic arborization in neuronal and brain slice cultures, whereas apoE4 inhibits these processes [21,22,37]. Neurite degeneration has been observed in transgenic mouse models of tauopathy [38] and in post-mortem AD patient brains [39], suggesting that dendrite abnormalities are an important component of AD-related neuropathology. Furthermore, it has been reported that neuron-derived apoE4 has a more detrimental effect on dendritic integrity compared to astrocytic apoE4 in response to excitotoxic injury, as evidenced by reduced MAP-2 immunostaining in NSE-apoE4, but not in GFAP-apoE4, transgenic mice [15]. In this study, we show that mice expressing apoE3 specifically in neurons have an increased number of total dendrites in both the CA1 and entorhinal cortex compared to neuronal apoE4–expressing mice, and the dendrites of NSE-apoE4 mice are shorter than those of NSE-apoE3 mice in these brain regions. We postulate that the toxicity of apoE4 produced by neurons contributes to the dendritic arborization impairments during aging and in response to brain injuries, while neuronal apoE3 promotes dendrite development.

In our experiments, neither apoE4-KI nor GFAP-apoE4 mice showed differences in dendritic arborization in the CA1 or entorhinal cortex compared to their respective apoE3 counterparts, while in the auditory cortex, GFAP-apoE4 mice had longer and more dendrites than GFAP-apoE3 mice. In contrast, previous
studies described a decrease in dendrite length and number of GFAP-apoE4 mice compared to GFAP-apoE3 mice [23]. These discrepancies might be due to several factors, including the age of the mice and the brain areas investigated. For instance, we examined the CA1 region of the hippocampus, the entorhinal cortex, and the auditory cortex at 7–8 and 19–21 months of age, while the previous study focused on the dentate gyrus of mice at 12 and 24 months of age [23]. Differences in spine labeling and...
Table 1. Summary of the effects of apoE genotype on dendritic spine morphology in the CA1 subregion of the hippocampus, the entorhinal cortex, and the auditory cortex in different apoE mice at 19–21 months of age.

| Spines Shape   | CA1     | Entorhinal Cortex | Auditory Cortex |
|----------------|---------|-------------------|-----------------|
|                | Total   | Apical | Basal | Total | Apical | Basal | Total | Apical | Basal |
| Stubby         | NSE-E4 < NSE-E3 | No effect | E4-KI < E3-KI | No effect | No effect | No effect | No effect | No effect | No effect |
| Thin           | NSE-E4 < NSE-E3 | NSE-E4 < NSE-E3 | E4-KI < E3-KI | No effect | No effect | No effect | No effect | No effect | No effect |
| Mushroom       | GFAP-E4 > GFAP-E3 | No Effect | E4-KI < E3-KI | GFAP-E4 > GFAP-E3 | No effect | NSE-E4 < NSE-E3 | NSE-E4 < NSE-E3 | GFAP-E4 > GFAP-E3 | GFAP-E4 > GFAP-E3 |
| Filopodia      | NSE-E4 < NSE-E3 | GFAP-E4 > GFAP-E3 | NSE-E4 < NSE-E3 | GFAP-E4 > GFAP-E3 | No effect | No effect | No effect | GFAP-E4 > GFAP-E3 | GFAP-E4 > GFAP-E3 |

For details, see data presented in Figures 7, 8, and 9.

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neurons and astrocytes might contribute to such detrimental effects.

There are a number of potential mechanisms through which neuronal apoE4 might exert its inhibitory effects on spine density and morphology. Neuronal apoE4 is susceptible to proteolytic cleavage, producing C-terminally truncated fragments that are associated with increased tau phosphorylation [3,4,13,14]. The accumulation of hyperphosphorylated tau in spines has been linked to disrupted synaptic function through the impairment of glutamate receptor trafficking and synaptic anchoring [43]. Thus, neuronal apoE4 might reduce spine numbers by enhancing tau phosphorylation and localization into spines. Astrocyte-derived apoE4, on the other hand, is not associated with apoE fragmentation and tau phosphorylation [13], which might explain...
why it does not inhibit spine density compared to astrocytic apoE3. Neuronal apoE4 and its fragments also impair mitochondrial integrity, dynamics, and function [3,12,44]. Normal mitochondrial dynamics and function are essential for the morphogenesis and plasticity of spines and synapses [45]. Recently, an inherited variable poly-T repeat genotype of mitochondrial protein TOMM40 has been suggested to be associated with the age-of-onset of AD and cognitive aging [28,46]. It has also been suggested that apoE4 interacts with TOMM40 to cause mitochondrial impairment and neurotoxicity [28]. In this regard, apoE4 or its fragments might interact with TOMM40 on the mitochondrial outer membrane, causing the release of cytochrome C from damaged mitochondria with subsequent apoptosis, which could be a therapeutic target for AD [28].

Figure 9. Quantification of spine subtypes in the auditory cortex (AC) in different apoE mouse models at 19–21 months of age. (A–H) Averaged densities of stubby (A–B), thin (C–D), mushroom (E–F), and filopodia (G–H) type protrusions in the auditory cortex for all genotypes. Values are mean ± SEM, n = 20–33 neurons from 3–5 mice for each group. *p<0.05, **p<0.01, ***p<0.001.
shows cellular source-dependent differences in its modulation of excitotoxic responses. Whereas both neuronal and astrocytic apoE3 protected dendrites and synapses against excitotoxic injury in the neocortex and hippocampus, apoE4 was excitoprotective only when produced by astrocytes [15]. The lack of this excitoprotective function of neuronal apoE4 is likewise due to its susceptibility to proteolysis [15].

In conclusion, our study demonstrates for the first time that the cellular source of apoE determines its effects on dendritic arborization, spine density, and spine morphology. Neuron-derived apoE4 has detrimental effects on dendritic arborization and spine development, whereas astrocytic apoE4 does not. These studies should further our understanding of the functional differences of apoE derived from different cellular sources in the brain and provide insights for drug development targeting apoE4's detrimental effects.

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Author Contributions
Conceived and designed the experiments: SJ YH. Performed the experiments: SJ SYY LL JK. Analyzed the data: SJ SYY LL JK YH. Wrote the paper: SJ YH.

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