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Diosgenin is an exogenous activator of 1,25D₃-MARRS/Pdia3/ERp57 and improves Alzheimer's disease pathologies in 5XFAD mice

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The aim of this study was to investigate the effects and the mechanism of diosgenin, a famous plant-derived steroidal sapogenin, on memory deficits in Alzheimer's disease (AD) model mice. Diosgenin-treated 5XFAD mice exhibited significantly improved performance of object recognition memory. Diosgenin treatment significantly reduced amyloid plaques and neurofibrillary tangles in the cerebral cortex and hippocampus. Degenerated axons and presynaptic terminals that were only observed in regions closely associated with amyloid plaques were significantly reduced by diosgenin treatment. The 1,25D₃-membrane-associated, rapid response steroid-binding protein (1,25D₃-MARRS) was shown to be a target of diosgenin. 1,25D₃-MARRS knockdown completely inhibited diosgenin-induced axonal growth in cortical neurons. Treatment with a neutralizing antibody against 1,25D₃-MARRS diminished the axonal regeneration effect of diosgenin in A β (1–42)-induced axonal atrophy. This is the first study to demonstrate that the exogenous stimulator diosgenin activates the 1,25D₃-MARRS pathway, which may be a very critical signaling target for anti-AD therapy.

e have hypothesized that the enhancement of brain function requires the reinforcement of neuronal networks, including neurite regeneration and synapse formation; therefore, we have been exploring the use of anti-Alzheimer's disease (AD) drugs in reconstructing neuronal networks in the damaged brain. In the last decade, several strategies for lowering Aβ have been studied in basic research and clinical trials as an alternative to enhancing cholinergic function. However, immunotherapy with bapineuzumab, a humanized anti-Aβ monoclonal antibody, did not improve cognitive function in a phase 2 trial¹. Although a γ-secretase inhibitor, LY450139, also reduced plasma and cerebrospinal fluid Aβ levels in humans, cognitive amelioration was not detectable². Neuritic atrophy and loss of synapses underlie the pathogenesis of AD and are located upstream of neuronal death in the Aβ cascade^{3,4}. The dysfunction of neurites and synapses is a direct cause of the memory deficit in AD. Because neurons with atrophic neurites may remain viable and have the potential to be remodeled, the essential event for the achievement of recovery of brain function after injury is the reconstruction of neuronal networks, including neurite regeneration and synaptic reformation⁵.

5XFAD mice are engineered to co-overexpress and co-inherit mutant human APP (the Swedish mutations: K670N and M671L; the Florida mutation: I716V; and the London mutation: V717I) and PS1 (M146L; L286V) transgenes under the neuron-specific mouse Thy-1 promoter⁶. Five familial AD mutations act together to additively increase the levels of cerebral A β peptides, especially the neurotoxic peptide A β 42. While the majority of AD transgenic mice require 6–12 months or longer to form amyloid plaques⁷, 5XFAD mice begin to develop visible amyloid deposits as early as 2 months of age, which is consistent with their dramatically accelerated A β 42 generation. In addition to the memory deficits in 5XFAD mice, demonstrated by contextual fear conditioning⁸ and their performance in a Y-maze⁶, we clarified the impairment of spatial memory⁹ and object recognition memory^{10,11} in these mice.

Previously, we found that the herbal drug-derived steroidal sapogenins regenerated neurite atrophy and synaptic loss, leading to memory improvement in AD model mice^{10,12}. Diosgenin is also a steroidal sapogenin

and a major constituent in Dioscorea rhizome and other herbal drugs, such as those from Trigonella spp., Polygonatum spp. and Smilax spp. Several biological effects of diosgenin have been reported, such as anti-cancer effects¹³, anti-food allergy effects¹⁴, anti-cognitive deficit effects¹⁵ and relief of diabetic neuropathy¹⁶. A diosgenin derivative, caprospinol (diosgenin 3-caproate), reduces amyloid deposits and improves memory dysfunction in AB1-42infused AD model rats¹⁷. This result led us to hypothesize that diosgenin might also improve memory impairment in 5XFAD mice by decreasing A_β. A variety of possible signaling pathways for diosgenin have been reported. For example, diosgenin enhanced PI3 kinase activity in melanogenesis¹⁸. In contrast, diosgenin attenuated the TNF-α-stimulated phosphorylation of Akt, ERK, JNK and p38 in a vascular smooth muscle cell line¹⁹. In hepatocellular carcinoma cell lines, diosgenin inhibited the phosphorylation of STAT3 and downstream c-Src, JAK1 and JAK2²⁰. However, the signaling mechanism of diosgenin in neuronal cells and the direct target protein remain unknown. Although several small-molecular-weight compounds derived from medicinal plants exhibit multiple bioactivities, the direct target proteins of those exogenous chemical compounds remain largely unknown. In the present study, we investigated the effects of diosgenin on memory deficits in 5XFAD mice and identified a direct target protein for diosgenin.

Results

To investigate the effects of diosgenin on impaired object recognition memory in 5XFAD mice, diosgenin (10 µmol/kg=4.14 mg/kg), memantine (200 µmol/kg=43.15 mg/kg) or vehicle solution was administered i.p. to mice for 20 days. The day after the last administration, an object recognition test was conducted. To assess the appropriate time interval between a training session and a test session, another group of mice had been tested previously, and the test session was conducted 10 and 30 min and 24 h after the training session. We had determined the appropriate time interval between a training session and a test session in which control 5XFAD mice cannot memorize an object to be 30 min. Wild-type mice and diosgenin-treated 5XFAD mice showed significantly more frequent exploratory behavior in the presence of a novel object than chance (50%) (Figure 1A). The value of the diosgenin-treated group in the test session was significantly higher than that in the training session. In contrast, the exploratory behavior in the presence of the novel object in the vehicle-treated 5XFAD mice was close to chance (50%). The memantine-treated group showed no improvement in memory. In the open field locomotion test, no significant differences were detected in the moving velocities and distances moved of the four groups in the test (data not shown).

The day after the novel recognition test, the brains of the mice were removed, and slices were prepared for immunohistochemistry. We measured the parietal area containing the parietal cortex and hippocampus at the bregma in 1.4–2-mm sections of the brain. The area of the amyloid deposits was measured in the brain slices. A large number of amyloid deposits were primarily localized in the cerebral cortex and hippocampus of 5XFAD mice. As shown in Figure 1B, the large A β (1–40/42)-positive stains (greater than 50 µm in width) were measured as extracellular amyloid deposits, although intracellular inclusions were also observed infrequently in the 5XFAD mice. In the age-matched wild-type mice, no amyloid plaque was observed. The diosgenin-treated 5XFAD mice exhibited a significant reduction of the amyloid plaques in the cerebral cortex and hippocampus (Figure 1C). Memantine also significantly reduced amyloid plaques in the cortex and hippocampus (Figure 1C).

Intense paired helical filament (PHF)-tau expression was observed very close to the amyloid plaques, as well as in the distal area of the amyloid plaques in the 5XFAD mice (Figure 1B). In the wild-type mice, no PHF-tau expression was observed. Diosgenin-treated 5XFAD mice showed a significant reduction of PHF-tau expression in the cortex (associated with and distal to the plaques) and the hippocampus (distal to the plaques) (Figure 1D). Memantine yielded no reduction of PHF-tau expression (Figure 1D).

Bulb-like axons, which were limited to the amyloid plaques, were markedly observed (Figure 2A). Abnormal axon structures, such as bulb shapes and dense staining by the pNF-H antibody, were reported in early- and end-stage AD patients³. In addition, in the PSAPP transgenic mouse AD model, abnormally swollen varicosities of neurites inside fibrillar amyloid deposits were shown in degenerated and dying axons²¹. With the exception of amyloid plaques in the brain, no abnormal structures were detected in the vehicle-treated 5XFAD and diosgenin-treated 5XFAD mice. The areas of bulb-like swollen axons in the cortex and hippocampus were significantly reduced by diosgenin treatment (Figure 2B). However, memantine treatment did not reduce the swollen axons (Figure 2B).

Abnormally swollen presynaptic boutons closely associated with amyloid plaques were also reported as degenerative alterations in 5XFAD mouse cortex²² and aged monkey cortex²³. Similar presynaptic alterations were observed in the current study. As shown in Figure 2A, synaptophysin-positive staining in the distal area of amyloid plaques exhibited a minute dot-like pattern in the 5XFAD mice. However, synaptophysin-positive staining associated with amyloid plaques in the vehicle-treated 5XFAD mouse was very dense and exhibited large, swollen dots. In contrast, the abnormal presynaptic structures were significantly reduced by diosgenin treatment in the cerebral cortex but not the hippocampus (Figure 2C). Memantine showed no effects on the reduction of presynaptic degeneration (Figure 2C).

We used the target identification using drug affinity responsive target stability (DARTS) method to identify a direct target protein for diosgenin in neurons²⁴. Cell lysate of mouse cortical neuron primary culture was added to diosgenin or vehicle solution and was then reacted with thermolysin. A 57-kDa protein band in the diosgenintreated lysate was shown to be thicker than in the vehicle-treated lysate on a silver-stained SDS-PAGE gel (Figure 3A). LC/MS/MS analysis indicated that the band was possibly the protein disulfideisomerase A3 (Pdia3). The Pdia3 protein has many synonyms, including ERp57, GRP58 and the membrane-associated rapid response steroid-binding receptor (1,25D₃-MARRS). This protein is present in multiple subcellular locations, including the endoplasmic reticulum, plasma membrane, cytosol and nucleus^{25,26}. 1,25D₃-MARRS was reported to be expressed on the cell surface and to mediate the rapid response of 1a,25-dyhydroxyvitamin D3 (DHVD3)²⁷. The genomic effects of DHVD3 are mediated by a nuclear receptor, nVDR, which operates through binding to DNA and activating gene expression. In contrast, the rapid and non-genomic cell response after the stimulation of DHVD3 is mediated by the cell surface function of 1,25D₃-MARRS²⁸ and activates numerous signal transduction cascades, including protein kinase C (PKC), protein kinase A (PKA), extracellular-response-activated kinase (ERK) and phosphoinositide 3-kinase (PI3K)^{27,29,30}. As the studies of 1,25D₃-MARRS have primarily used intestinal and osteoblast cells, its function in the brain has never been investigated. The conformation state of DHVD3 is 6-s-trans, which causes it to associate mainly with nVDR³¹. In contrast, the 6-s-cis conformation of DHVD3 and a 6-s-cis locked derivative of DHVD3, 1a,25-dyhydroxylumisterol, stimulated a rapid response³². Because the core structure of diosgenin is similar to that of the derivative 1a,25-dyhydroxylumisterol, we focused on the possibility that 1,25D3-MARRS might mediate the diosgenin signal as a cell surface receptor.

The expression of $1,25D_3$ -MARRS was investigated in cortical cell primary culture using a specific antibody for the $1,25D_3$ -MARRS, Ab099 clone. An Ab099-positive signal was distributed mainly in neurons but not astrocytes (Figure 3B). A cell binding assay was performed to detect the binding activity of diosgenin to cell-surface $1,25D_3$ -MARRS. Without triton-X, the binding of DHVD3 on

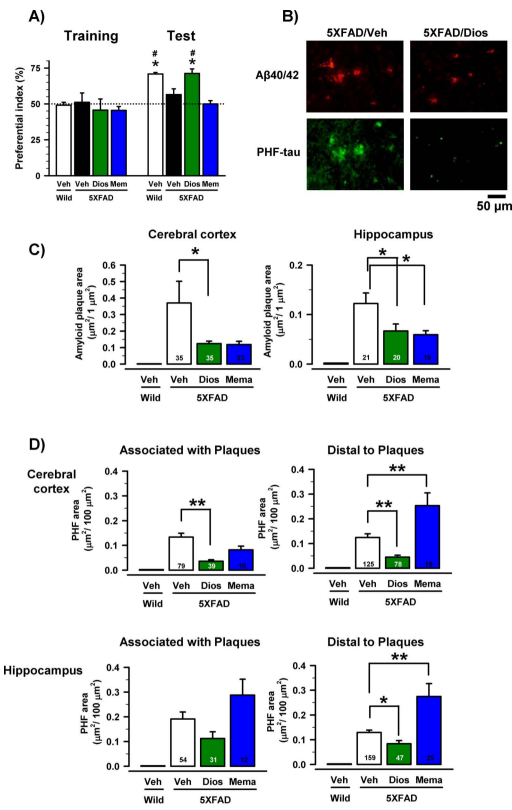
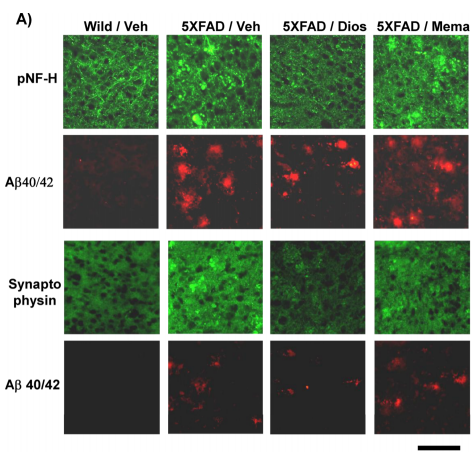


Figure 1 | Effects of diosgenin on object recognition memory deficits and AD pathologies in 5XFAD mice. Diosgenin (10 μ mol/kg, i.p.), memantine (200 μ mol/kg, i.p.) or vehicle solution was administered for 20 days to mice (females, 6–8 months old). The day after the last administration, an object recognition test was carried out. (A) The preferential indices of the training and test sessions are shown. (*p < 0.05 vs. vehicle-treated 5XFAD mice, one-way ANOVA *post hoc* Dunnett's test; *p < 0.05 vs. the same group in the training session, paired *t*-test; n=4–6 mice). (B–D) The total area of amyloid plaques and PHF-tau in the cerebral cortex and hippocampus were measured in vehicle-treated wild-type, vehicle-treated, diosgenin-treated and memantine-treated 5XFAD mice. (B) The photographs are representative images of A β (1–40/42)-positive plaques and PHF-tau in the cerebral cortex. (C) The total area of amyloid plaques per 1 μ m² was quantified in the cerebral cortex and hippocampus. The regions associated with amyloid plaques and distal to amyloid plaques were measured separately (*p < 0.05, **p < 0.001, one-way ANOVA *post hoc* Dunnett's test; the number of measured areas is shown in each column).





100 μm

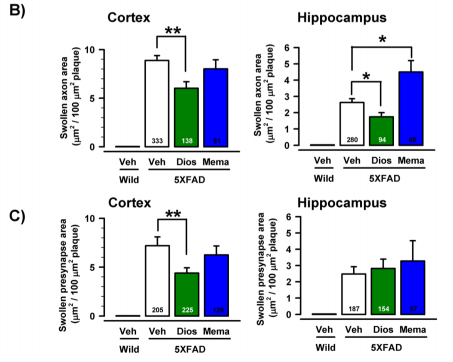


Figure 2 | Effects of diosgenin on axonal and presynaptic abnormalities associated with amyloid plaques. (A) Double labeling with antibodies to $A\beta(1-40/42)$ (red) and pNF-H (green) or $A\beta(1-40/42)$ (red) and synaptophysin (green) was carried out. Bulb-like axonal structures positive for pNF-H antibody and swollen puncta positive for synaptophysin antibody were localized only in the amyloid plaque areas. Representative images of the cerebral cortex of vehicle-treated wild-type (Wild/Veh) mice, as well as 5XFAD (5XFAD/Veh), diosgenin-treated 5XFAD (5XFAD/Dios) and memantine-treated 5XFAD (5XFAD/Mema) mice, are shown. (B) The quantified areas of abnormal axons per 100 μ m² of amyloid plaque are shown in the cerebral cortex and hippocampus. (C) The quantified areas of abnormal presynapses per 100 μ m² of amyloid plaque are shown in the cerebral cortex and hippocampus (*p < 0.05, **p < 0.001, one-way ANOVA *post hoc* Dunnett's test; the number of measured areas is shown in each column).



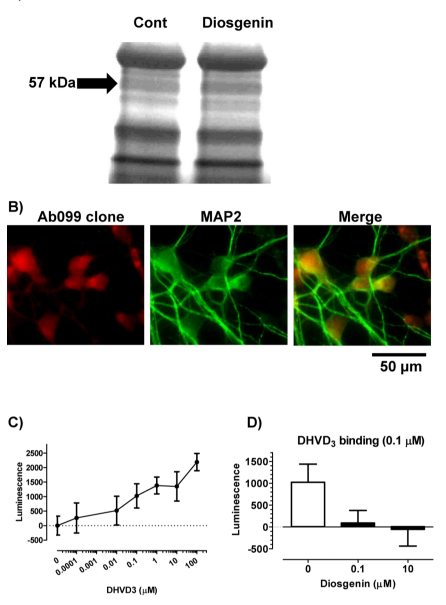


Figure 3 | 1,25D₃-MARRS as a direct target protein of diosgenin. (A) Mouse cortical neuron primary cultures (ddY, E14) were maintained for 24 h without drug treatment. Cell lysate was added to diosgenin or vehicle solution and incubated for 30 min at 4°C. The mixture was proteolysed using thermolysin and electrophoresed. The separated proteins were stained. The band in the diosgenin-treated lysate (diosgenin lane) that was thicker than that in the vehicle-treated lysate (Cont lane) was cut out and prepared for mass spectrometry analysis. (B) Double labeling with antibodies to 1,25D₃-MARRS (Ab099 clone, red) and MAP2 (green) was carried out in primary cultures of rat cortical neurons. (C) The dose-dependency of cell surface binding of DHVD3 in primary culture cortical neurons. Diosgenin (0–100 μ M) was added to cells and incubated for 10 min. The DHVD3 binding to the cell-surface was detected using immunocytochemistry with an anti-DHVD3 antibody. (D) The replacement of the cell-surface binding of DHVD3 binding to the cell surface was detected by immunocytochemistry using an anti-DHVD3 antibody.

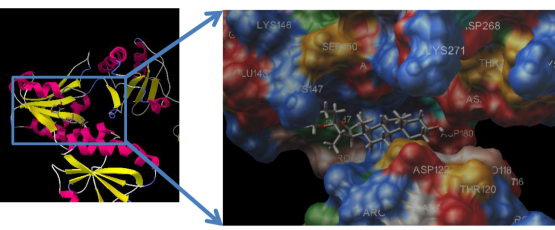
cortical neurons was detected using an immunoluminescence assay. DHVD3 binding was detected in a dose-dependent manner (0.0001–100 μ M) (Figure 3C). Because 0.1 μ M DHVD3 exhibited submaximal binding, the replacement of DHVD3 (0.1 μ M) with diosgenin was investigated. The simultaneous treatment with diosgenin (0.1 and 10 μ M) reduced the DHVD3 (0.1 μ M) binding (Figure 3D). The binding sites of 1,25D₃-MARRS and nVDR to DHVD3 and diosgenin were predicted using the docking suite Autodock 4.0 (http://autodock.scripps.edu/). Figure 4 shows the calculated binding structure of 1,25D₃-MARRS to diosgenin. The groove area between the α and β domains of 1,25D₃-MARRS contacts diosgenin (Figure 4). The docking scores of diosgenin and DHVD3 to

 $1,25D_3$ -MARRS were -8.4 and -7.9, respectively (Table 1), and the docking scores of diosgenin and DHVD3 to nVDR were -8.2 and -12.6, respectively (Table 1). These results suggest that the binding potency of diosgenin to $1,25D_3$ -MARRS is slightly higher than that of DHVD3 and that the binding potency of DHVD3 to nVDR is much higher than that of diosgenin.

To investigate the function of 1,25D₃-MARRS in diosgenininduced axonal extension, 1,25D₃-MARRS was knocked down using siRNA transfection. Two days after the transfection of siRNA for 1,25D₃-MARRS (400 nM), the 1,25D₃-MARRS level was markedly reduced in cortical neurons (Figure 5A). In contrast, nVDR expression in cortical neurons was not changed by the transfection of siRNA

A)





B)

Figure 4 | Docking simulation of diosgenin to 1,25D₃-MARRS. The predicted binding complex site of 1,25D₃-MARRS and diosgenin is shown (A). The groove area between the α domain and the β domain of 1,25D₃-MARRS contacts diosgenin (B).

for 1,25D₃-MARRS, suggesting that the knockdown did not affect the genomic signaling of DHVD3 (Figure 5A). Therefore, 2 days after siRNA transfection, denosomin (1 μ M), DHVD3 (1 μ M) or vehicle solution was applied to the cells. After incubation for 4 additional days, the cells were double-immunostained for pNF-H and MAP2 to measure the axonal length per neuron. In cortical neurons, after the transfection of control siRNA, the density of axons was significantly increased by diosgenin and DHVD3 treatments (Figure 5B). However, both diosgenin-induced and DHVD3-induced axonal growth were almost completely inhibited in neurons after the transfection of 1,25D₃-MARRS siRNA (Figure 5B). These data indicate that 1,25D₃-MARRS is essential for the diosgenin-induced extensions of axons.

A)

To investigate which protein kinases are involved in diosgenininduced axonal growth signaling, a specific inhibitors for PI3K, MEK1, PKC or PKA was co-applied with diosgenin. The doses of inhibitors used were determined from the known IC₅₀ values. The PI3K inhibitor LY294002 (10 μ M), the MEK1 inhibitor PD98059 (1 and 10 μ M), the protein kinase C inhibitor Gő6970 (10 and 100 nM) and the protein kinase A inhibitor PKI (0.1 and 1 μ M) completely inhibited diosgenin-induced axonal growth (Figure 6A). DHVD3-induced axonal growth was also completely inhibited by LY294002 (1 and 10 μ M), PD98059 (1 and 10 μ M), Gő6970 (100 nM) and PKI (0.1 and 1 μ M).

We investigated the effects of diosgenin on axonal regeneration *in vitro* and the effects of neutralizing 1,25D₃-MARRS on diosgenininduced axon regeneration in rat cortical neurons. Three days after treatment with A β (1–42), normal IgG as a control or Ab099, a specific neutralizing antibody for 1,25D₃-MARRS²⁸ was applied to the cells. After the 10-min incubation, cells were treated with diosgenin (0.1 and 1 μ M) or vehicle solution. A β (1–42) treatment markedly reduced the density of axons. In contrast, treatment with Ab099 completely diminished the axonal growth induced by diosgenin (Figures 7A and 7B).

Table 1 Docking scores of ligands and receptors		
	Diosgenin	DHVD3
1,25D ₃ -MARRS nVDR	-8.4 -8.2	-7.9 -12.6

Discussion

The 5XFAD mouse model reveals AD pathologies at an early age. Memory dysfunctions in the 5XFAD mice, including deficits in object recognition memory^{10,11}, spatial memory⁹ and contextual fear memory⁸, were detected at 4 months old. The present study showed for the first time that diosgenin administration improved the object recognition memory deficit in 5XFAD mice. Diosgenin administration inhibited several signs of neuronal degeneration, including presynaptic degeneration associated with amyloid plaques in the cortex, axonal degeneration associated with amyloid plaques in the cortex and hippocampus and PHF-tau expression associated with and distal to amyloid plaques in the cortex and hippocampus. In addition, amyloid plaques were decreased by diosgenin treatment. It was reported that the diosgenin derivative, caprospinol (diosgenin 3caproate), reduced amyloid deposits and improved memory dysfunction in Aβ1-42-infused AD model rats¹⁷. Therefore, it is possible that the amyloid plaque decreasing effect of diosgenin results in the protection of axonal degeneration and the hyperphosphorylation of tau. However, diosgenin can enhance axonal growth in normal neurons (Figures 5 and 6) and can induce the regrowth of axons in Aβtreated neurons following treatment (Figure 7). In addition, diosgenin shows no direct binding activity to $A\beta^{33}$. These results indicate that diosgenin may contribute to axonal extension through a direct pathway, as well as through the amyloid plaque-lowering pathway. The approved anti-AD drug memantine caused no improvement in object recognition (Figure 1B), the degeneration of axons and presynapses (Figure 1D) or PHF-tau expression (Figures 2B and 2C), but it did reduce amyloid plaques (Figure 1C). The reduction of amyloid plaques by memantine was previously reported in the AD model Tg2576 mice at 10 and 20 mg/kg34 and APP/PS1 mice at 10 mg/kg³⁵, although the mechanism by which it occurred is unknown. Conflicting results have been reported concerning memory improvement due to memantine. Fear-conditioned memory is not improved by treatment with 10 or 20 mg/kg memantine for 6 months in Tg2576 mice, despite the reduction of amyloid plaques³⁴. The administration of 10 mg/kg memantine for 4 months improved object recognition in APP/PS1 mice³⁵, but the administration of 10 mg/kg memantine for 1 week did not affect the spatial memory deficit in APP23 mice³⁶. Surprisingly, memantine treatment significantly increased the number of degenerated axons in the dentate gyrus of Tg2576 mice³⁴. Additionally, in our study, memantine increased the number of degenerated axons in the hippocampus (Figure 2). These results suggest that memantine may have both neuroprotective and neurotoxic effects. In our previous study, a



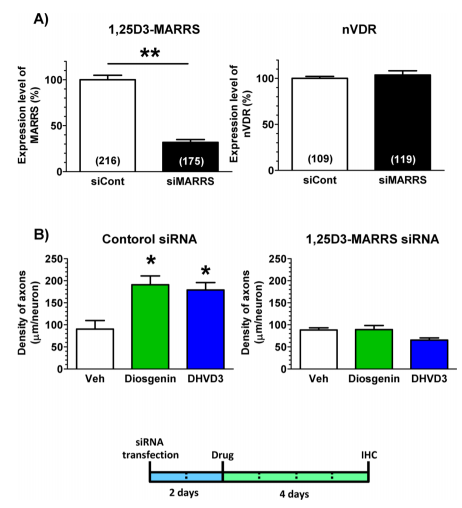


Figure 5 | The effect of 1,25D₃-MARRS knockdown on diosgenin-induced axonal outgrowth. (A) siRNA for Pdia3 (400 nM) or control siRNA (400 nM) was transfected into rat cortical neurons. Two days later, double labeling with antibodies to 1,25D₃-MARRS (Ab099 clone) and MAP2, or to nVDR and MAP2, was carried out. The expression levels of 1,25D₃-MARRS and nVDR in neurons were quantified (*p < 0.05, *p < 0.001, one-way ANOVA *post hoc* Dunnett's test; the number of measured areas is shown in each column). (B) siRNA for Pdia3 (400 nM) or control siRNA (400 nM) was transfected into cortical cells. Two days later, the cells were incubated with 1 μ M diosgenin, 1 μ M DHVD3 or the vehicle solution (Veh, 0.1% DMSO). Four days after drug administration, the cells were fixed and double-immunostained for pNF-H and MAP2. The density of pNF-H-positive axons per MAP2-positive neuron was quantified for each treatment (*p < 0.05, one-way ANOVA *post hoc* Dunnett's test, n=12–14).

higher dose of diosgenin (100 μ mol/kg) was administered to 5XFAD mice (male, 7–9 months old) by i.p. injection for 19 days. In object recognition test, diosgenin-treated 5XFAD mice showed memory improvement. Since the efficacy of diosgenin at a dose of 100 μ mol/kg was almost same to that of 10 μ mol/kg diosgenin, we consider that 10 μ mol/kg diosgenin is enough to improve memory dysfunction.

Figure 2 shows that drastic axonal and presynaptic morphological changes occurred and were associated with amyloid plaques in 5XFAD mice. Other groups have also reported these axonal and synaptic degenerations in AD patients³, 5XFAD mice²² and other AD model mice²¹. Because a bulb-shaped axon overlaps with a swollen presynapse in the early and late stages of AD³⁷, a terminal of the degenerated axon may also be degenerated, and synaptophysin-positive synaptic vesicles may be diffused. The core structure of degenerated axons and presynapses is called a neuritic plaque, which is one of the pathological hallmarks of AD. These morphological degenerations of axons and presynapses may lead to the disruption of neuronal networks. It is possible that A β accumulation is closely related to the dystrophy of neurites and synapses. For example, the application of an anti-A β antibody to the exposed brain surface of the PDAPP mouse, an AD model, reduced the number and size of

swollen dystrophic neurites around the site of amyloid plaques³⁸. In addition, axonal dystrophy triggers the production and release of Aβ, leading to the progression of neuritic plaque formation²². Axonal dystrophy is recognized as a critical degeneration associated with amyloid plaques. Because diosgenin-induced reductions of degenerated axons and presynapses were observed in each amyloid plaque, these reductions may be independent of the decrease in amyloid plaques caused by diosgenin. Memantine reduced amyloid plaques but did not provide amelioration of axonal and presynaptic degeneration, also suggesting that decreasing the number of amyloid plaques is not necessary or sufficient for normalizing axonal structure and function. We then investigated the diosgenin signaling pathway to clarify the mechanism of restoration of axonal degeneration in AD.

Interestingly, our results indicated that the receptor mediating DHVD3 signaling also works as a diosgenin receptor. Although the function of 1,25D₃-MARRS in neurons and neurodegenerative diseases was not unknown, the receptor, as a rapid and non-genomic response mediator, was considered to interact with multiple signaling pathways in intestinal and osteoblast cells, including PKA^{27,28}, PKC^{27,29,39}, ERK²⁹, PI3K³⁰ and STAT3²⁶. Our results also show that diosgenin stimulates PI3K, ERK, PKC and PKA (Figure 6). However, no reports have yet elucidated why the multiple signaling pathways

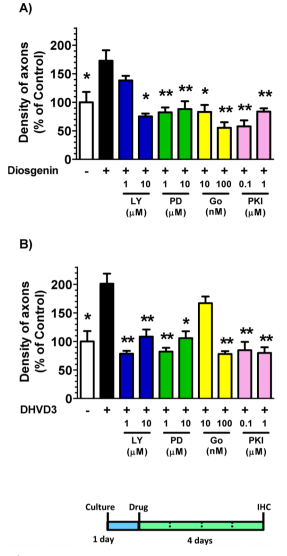


Figure 6 | Effects of protein kinase inhibitors on diosgenin-induced axonal growth. Cortical neurons were cultured for one day and then treated with 1 μ M diosgenin (A), DHVD3 (B) or vehicle solution. Simultaneously, the PI3K inhibitor LY294002 (10 μ M), the MEK1 inhibitor PD98059 (1 and 10 μ M), the protein kinase C inhibitor Gő6970 (10 and 100 nM) or the protein kinase A inhibitor PKI (0.1 and 1 μ M) was applied to the cells. Four days after the treatment, the cells were fixed and double-immunostained for pNF-H and MAP2. The density of pNF-H-positive axons per MAP2-positive neuron was quantified for each treatment (*p < 0.05, **p < 0.001, one-way ANOVA *post hoc* Dunnett's test, n=6–7).

can be activated by 1,25D₃-MARRS. It also remains unknown how 1,25D₃-MARRS is associated with the plasma membrane, although 1,25D₃-MARRS has one myristoylated site at amino acid residues 315–320⁴⁰. Although a neurite outgrowth effect of DHVD3 was reported in embryonic rat hippocampal neurons, it is unknown whether the effect is mediated by a genomic or non-genomic response⁴¹. The PI3K-Akt pathway is well known to regulate local protein translation via the mTOR pathway, thus playing an important role in axon regeneration⁴². PI3K also regulates Cdc42, which is a key regulator of cytoskeletal reorganization in axonal tips⁴³. ERK signaling is also required for local axon assembly⁴⁴ and local protein translation at the growth cone⁴⁵. The phosphorylation of GAP-43 by PKC in growth cones is required for axonal outgrowth⁴⁶. The PKA pathway is also known to be associated with axonal extension in cortical neurons⁴⁷. The non-genomic action of DHVD3 is known to enhance

the cerebral clearance of the A β (1–40) peptide in mice⁴⁸, and ERp57 is one of the carrier proteins that prevents the aggregation of $A\beta$ in cerebrospinal fluid⁴⁹, suggesting that the diosgenin-stimulated 1,25D₃-MARRS pathway may contribute to AB elimination from the brain. The PI3K-Akt and MEK-ERK pathways are reported to promote nonamyloidogenic cleavage of APP, resulting in a decrease in A β^{50} . Although we need to investigate other possible causes of A β lowering by diosgenin, such as β -secretase, γ -secretase, A β accumulation inhibitors and Aβ-degrading enzymes, the multiple signaling pathways stimulated by diosgenin may contribute to the enhancement of axonal growth and Aß reduction. Our unpublished study for identifying phosphorelated proteins by diosgenin using 5XFAD mice showed that downstream molecules of PI3K activation (PDK1, Akt1 and XIAP), a PKA substrate (CREB), a PKC substrate (MARCKS) and upstream molecules of MEK1 (Raf1 and MEKK1) and a downstream molecule (RSK2) were phosphorylated 30 min after diosgenin treatment in the hippocampus (data not shown). Those results suggest that diosgenin may stimulate multiple signal pathways also in vivo.

The nVDR-mediated genomic action of DHVD3 against axonal damage has not been elucidated, although nVDR expression is rich in most areas of the brain⁵¹. It has been reported that the *in vitro* exposure of cortical neurons to A β 1-42 reduced nVDR expression and induced cytotoxicity, and the addition of DHVD3 up-regulated nVDR expression and inhibited cell death⁵². However, there are no reports demonstrating any relation between axonal degeneration and nVDR. In our study, nVDR was not involved in diosgenin- or DHVD3-induced axonal growth in cortical neurons (Figure 5), suggesting that the 1,25D₃-MARRS cascade is related to axonal growth and regrowth events.

A docking simulation suggested that the binding potency of diosgenin to 1,25D₃-MARRS is greater than that of DHVD3. In contrast, the binding potency of diosgenin to nVDR is less than that of DHVD3. Although the conformation of DHVD3 is flexible, approximately 95% of it exists in the non-steroidal and extended 6-s-trans form^{31,53}. In addition, the biological concentration of DHVD3 is strictly controlled and metabolized to 1 α ,24,25-dihydroxyvitamin D3⁵⁴ because hyper DHVD3 causes adverse effects, such as hypercalcemia. Therefore, diosgenin is an exogenous stimulator that allows the 1,25D₃-MARRS cascade to work more effectively than the endogenous agonist.

This is the first report of diosgenin successfully recovering the memory deficit in 5XFAD mice and restoring axonal and presynaptic degeneration in the cerebral cortex and hippocampus. We also found that diosgenin may function as an exogenous stimulator of $1,25D_3$ -MARRS and induce axonal growth and regrowth, even under A β -induced damaging conditions. These results suggest that the $1,25D_3$ -MARRS pathway is activated by the exogenous stimulator diosgenin and may be a very critical signaling pathway for anti-AD therapy.

Methods

All experiments were performed in accordance with the Guidelines for the Care and Use of Laboratory Animals of the Sugitani Campus of the University of Toyama. All protocols were approved by the Committee for Animal Care and Use of the Sugitani Campus of the University of Toyama. The approval number for the animal experiments is S2008-INM-6, and the confirmation number for the gene recombination experiments is 20-1. All efforts were made to minimize the number of animals used.

Materials. Diosgenin (Wako, Osaka, Japan), memantine hydrochloride (Tokyo Chemical Industry, Tokyo, Japan) and 1 α ,25-dihydroxy vitamin D₃ (DHVD3), Cayman Chemicals, Ann Arbor, MI, USA) were dissolved in dimethyl sulfoxide (DMSO). A β (1–42) (Sigma-Aldrich, St. Louis, MO, USA) was dissolved in sterile distilled water and incubated for 4 days at 37°C. The phosphatidylinositol 3-kinase (P13K) inhibitor LY294002 (Cayman Chemicals), MEK1 inhibitor PD98059 (LC laboratories) and protein kinase C inhibitor Gő6970 (LC laboratories, Woburn, MA, USA) were dissolved in DMSO. The protein kinase A inhibitor PKI (14–22) amide (myristoylated) (Enzo, Farmingdale, NY, USA) was dissolved in distilled water.

Animals. Transgenic mice (5XFAD) were obtained from the Jackson Laboratory (Bar Harbor, ME, USA). The 5XFAD mice have the following five mutations: Swedish



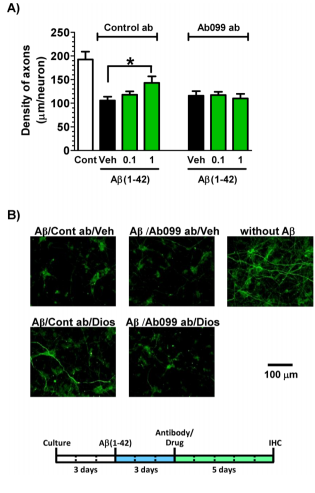


Figure 7 | Effect of the 1,25D₃-MARRS neutralizing antibody on diosgenin-induced axonal regeneration. Cortical neurons were cultured for three days and then treated with or without aggregated A β (1–42) (5 μ M). Three days after the administration of A β (1–42), the cells were treated with Ab099 antibody (Ab099 ab) or normal rabbit IgG (Control ab). After a ten-minute incubation period, diosgenin (0.1 and 1 μ M) or vehicle solution (0.1% DMSO, Veh) was administered to the cells. Five days after treatment, the cells were fixed and double-immunostained for pNF-H and MAP2. The density of pNF-H-positive axons per MAP2-positive neuron was quantified for each treatment (A) (*p < 0.05, one-way ANOVA *post hoc* Dunnett's test, n=7–9). (B) Representative images of each treatment are shown. The concentration of diosgenin was 1 μ M.

(K670N and M671L), Florida (I716V) and London (V717I) in human APP695 cDNA and human PS1 cDNA (M146L and L286V) under the transcriptional control of the neuron-specific mouse Thy-1 promoter⁶. They were maintained by crossing hemizygous transgenic mice with B6/SJL F1 breeders. To investigate the effect of diosgenin on 5XFAD, the study used hemizygous 5XFAD mice (female, 6–8 months old) and non-transgenic littermate wild-type mice (female, 6–8 months old), which were obtained by crossing a hemizygous 5XFAD mouse and a B6/SJL F1 mouse. All mice were housed with free access to food and water and were kept in a controlled environment ($22 \pm 2^{\circ}$ C, $50 \pm 5\%$ humidity, 12-h light/dark cycle starting at 7:00 am).

Novel object recognition test. Diosgenin or memantine hydrochloride was dissolved in DMSO at 10 times the final concentration, and the stock solution was diluted in physiological saline to the final concentration immediately before treatment. The drug or vehicle solution (10% DMSO in physiological saline) was intraperitoneally administered once a day for 20 days. On the last day of drug administration, mice were individually habituated to an open-field box composed of polyvinyl chloride (33 cm×28 cm; height, 26.5 cm) for 10 min. Their paths were tracked using a digital camera system. The distance moved for 10 min was analyzed as the locomotion activity with EthoVision 3.0 (Noldus, Wageningen, The Netherlands). The next day, a novel object recognition test was performed as described previously^{10,11}. Testing was carried out in a dimly illuminated room. Immunohistochemistry. One day after the novel object recognition test, mice were anesthetized and transcardially perfused with cold physiological saline. The brains were carefully removed from the skull, immediately immersed in 10 to 30% sucrose-PBS and stored at -80° C. The brains were cut in 20-µm successive coronal slices every 100 µm in the parietal area (bregma 1.4-2 mm) sections using a cryostat (CM3050S, Leica, Heidelberg, Germany). The slices were fixed with 4% paraformaldehyde and stained with a polyclonal antibody against $A\beta(1-40/42)$ (1:300) (Chemicon, Temecula, CA, USA), a monoclonal antibody against pNF-H (1:500) (Covance, Emeryville, CA, USA), a monoclonal antibody against synaptophysin (1:500) (Sigma-Aldrich) and a monoclonal antibody against PHF-tau (1:100) (Thermo Scientific, Rockford, IL, USA) at 4°C for 20 h. Alexa Fluor 488conjugated goat anti-mouse IgG (1:300) and Alexa Fluor 568-conjugated goat antirabbit antibody (1:300) were used as secondary antibodies (Molecular Probes Eugene, OR, USA). The fluorescent images for axons, presynapses, PHF-tau and $A\beta(1-40/42)$ were captured using a fluorescent microscope (AX-80) at 324 μ m \times 430 μ m (for pNF-H, synaptophysin and PHF-tau) or 1620 μ m \times 2150 μ m (for A β (1-40/42)). Three successive brain slices of the frontal cortex and five successive slices of the hippocampus were captured from a mouse for quantification. Extracellular amyloid plaques were determined by the size (greater than 50 µm in width). The area of amyloid plaques was measured using the image analyzing software ImageJ (http://rsbweb.nih.gov/ij). The area of the pNF-H-positive abnormal bulb-like axons was measured using ImageJ for every amyloid plaque in the fluorescent images (324 μ m×430 μ m). The areas of the synaptophysin-positive presynaptic boutons in every amyloid plaque were measured in fluorescent images $(324 \,\mu\text{m} \times 430 \,\mu\text{m})$ using Image J. The areas of PHF-tau in every amyloid plaque and outer amyloid plaques were measured in fluorescent images (324 µm×430 µm) using ImageI.

Primary culture. Embryos were removed from pregnant Sprague-Dawley rats (Japan SLC, Shizuoka, Japan) at 17 days of gestation. The cortices were dissected, and the dura mater was removed. The tissues were minced, dissociated and grown in cultures with neurobasal medium (Invitrogen, Grand Island, NY, USA) that included 12% B-27 supplement (Invitrogen), 0.6% D-glucose and 2 mM L-glutamine on 8-well chamber slides (Falcon, Franklin Lakes, NJ, USA) coated with 5 μ g/ml poly-D-lysine at 37°C in a humidified incubator with 10% CO₂. The seeding cell density was 4.35×10⁴ cells/cm².

Measurement of axonal density. To measure the density of axons, the cells were treated with 5 μ M A β (1-42) for 3 days (Figure 7) or without A β (1-42) for 4 days (Figures 5 and 6), and they were treated with diosgenin or vehicle (0.1% DMSO) for 3 days. The A β (1-42) was previously incubated for 4 days at 37°C for aggregation. When treated with the neutralizing antibody, the control antibody (normal rabbit IgG, 1:5000) or the polyclonal anti-rabbit 1,25D3-MARRS (Ab099 clone, 1:500) was applied to the cells 3 days after the A $\beta(1-42)$ treatment. Ten minutes after incubation with the antibody, the cells were treated with vehicle solution (0.1% DMSO) or diosgenin (0.1 and 1 µM). The cells were fixed with 4% paraformaldehyde and immunostained at 4°C for 20 h with a monoclonal antibody against pNF-H (1:500) as an axonal marker and a polyclonal antibody against MAP2 (1:500) as a neuronal marker. Alexa Fluor 488-conjugated goat anti-mouse IgG (1:300) and Alexa Fluor 568-conjugated goat anti-rabbit IgG (1:300) were used as secondary antibodies. The fluorescent images were captured using a fluorescence microscope system (BX61/ DP70, Olympus) at 324 µm×430 µm. Twelve to fourteen images (Figure 5B), 6 to 7 images (Figure 6) or 7 to 9 images (Figure 7A) were captured per treatment. The lengths of the pNF-H-positive axons were measured using an image analyzer Neurocyte (Kurabo), which automatically traces and measures neurite length without measuring the cell bodies. The sum of the axon lengths was divided by the number of MAP2-positive neurons. The resulting axon densities were averaged over all the images and are shown in Figures 5B, 6 and 7A.

Identification of target proteins for diosgenin. A mouse cortical neuron primary culture (ddY, E14) was maintained for 24 h without drug treatment. The cells were lysed with M-PER (Thermo Scientific) supplemented with protease and phosphatase inhibitors. After centrifugation (20,000 x g) for 15 min, the protein concentration of the lysates was determined using Quick Start (Bio-Rad Laboratories, Hercules, CA, USA). The cell lysate (5.6 μ g each) was added to 2 μ l of 10 mM diosgenin of vehicle solution and incubated for 30 min at 4°C. The mixture was proteolyzed using 7.5 μ g thermolysin in reaction buffer [50 mM Tris·HCl (pH 8.0), 50 mM NaCl, 10 mM CaCl₂] for 10 min at room temperature. To stop the proteolysis, 0.5 M EDTA (pH 8.0) was added to each sample in a 1:10 ratio. The samples were then mixed well and placed on ice.

The reaction samples were electrophoresed using a 5–20% real gel plate (Bio Craft, Tokyo, Japan). The separated proteins were stained using a negative gel stain MS kit (Wako). The bands in the diosgenin-treated lysate that were thicker than those in the vehicle-treated lysate were cut out and prepared for mass spectrometry analysis. After alkylation using iodpacetamide and in-gel digestion by trypsin, the supernatants were analyzed using nano-LC/MS/MS on a ThermoFisher LTQ Orbitrap XL. The MS/MS data were searched using Mascot (www.matrixscience.com).

Luminescence detection of 1,25D₃-MARRS binding. Primary cultured cortical neurons (SD, E17) were harvested in a 96-well white plate at a density of 0.5×10^5 cells/well. Three days later, the cells were treated with DHVD3 alone or in combination with DHVD3 and diosgenin for 10 min at 37°C. The cells were fixed

with 4% paraformaldehyde and incubated at 4°C for 20 h with a monoclonal antibody against DHVD3 (1:2000, Acris Antibodies, Herford, Germany) in Triton-X-free PBS to detect only the cell-surface binding of DHVD3. Horse radish peroxidase-conjugated goat anti-mouse IgG (1:2000) was used as a secondary antibody. The chemiluminescence of the antigen-antibody complex was detected using a GENios multi-plate reader (Tecan, Männedorf, Zurich, Switzerland).

siRNA transfection. siRNAs were transfected into rat cortical neurons (SD E18) according to the manufacturer's protocol for nucleofection (Lonza, Bazel, Switzerland). In brief, rat cortical neurons (5.25×10^6 cells) were mixed with 400 nM siPdia3 (a mixture of three sequences of Stealth siRNA for Pdia3, Invitrogen) or 400 nM control siRNA (Stealth RNAi Negative Control Low GC Duplex #3, Invitrogen) and electroporated with Amaxa Nucleofector (Lonza). Two days after the transfection, cells were fixed and double-immunostained with a monoclonal antibody against MAP2a and 2b (1:500, NeoMarker, Fremont, CA, USA), as a neuron marker, and Ab099 clone, a specific antibody for 1,25D₃-MARRS (1:500, provided as a gift by Dr. Nemere). Rat polyclonal anti-nVDR antibody (1:200, Millipore, Billerica, MA, USA) was used for the detection of the nuclear receptor for vitamin D3 (nVDR). The appropriate concentration of siRNA and the appropriate duration for knockdown (2 days after transfection) were determined previously. Two days after the transfection of siRNA, the cells were treated with diosgenin (1 μ M), DHVD3 (1 μ M) or vehicle solution (0.1% DMSO). Four days after treatment, the axon density was measured.

Docking simulation. Autodock 4.0 (AutoDock4 and AutoDockTools4; Molecular Graphics Laboratory. Department of Molecular Biology The Scripps Research Institute; http://autodock.scripps.edu/) was used to predict the conformation of the protein-ligand complexes⁵⁵. Ligand structures (diosgenin and DHVD3) were constructed and minimized using MOPAC2009 for 2D and 3D conformation⁵⁶. For the docking studies, the crystal structures of the proteins (PDB ID: NP_036630.1 for RLAR; NP_001619.1 for human AR) were determined from the protein sequence alignment [Brookhaven Protein Data Bank (PDB ID: 3f8u chain A for 1,25D₃-MARRS; 1ie9 chain A for nVDR)]. The predicted protein ligand complexes were optimized and ranked according to the empirical scoring function, which estimates the binding free energy of the ligand receptor complex.

Statistical analysis. Statistical comparisons were performed using a one-way analysis of variance (ANOVA) *post hoc* Dunnett's test and a paired *t*-test using SigmaStat 3.5 (SYSTAT, Chicago, IL, USA) and Graphpad Prism 5 (Graphpad Software, La Jolla, CA, USA). Values of p < 0.05 were considered significant. The means of the data are presented together with the SE.

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Author contributions

C.T. designed the study, wrote the protocol, carried out experiments and statistical analyses and prepared the manuscript. T.U. and M.U. carried out experiments and analyses. I.N. prepared and gifted an antibody. T.K. managed the literature searches and analyses. All authors contributed to and have approved the final manuscript.

Additional information

Competing financial interests: The authors declare no competing financial interests.

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