MAP2 is required for dendrite elongation, PKA anchoring in dendrites, and proper PKA signal transduction

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Microtubule-associated protein 2 (MAP2) is a major component of cross-bridges between microtubules in dendrites, and is known to stabilize microtubules. MAP2 also has a binding domain for the regulatory subunit II of cAMP-dependent protein kinase (PKA). We found that there is reduction in microtubule density in dendrites and a reduction of dendritic length in MAP2-deficient mice. Moreover, there is a significant reduction of various subunits of PKA in dendrites and total amounts of various PKA subunits in hippocampal tissue and cultured neurons. In MAP2-deficient cultured neurons, the induction rate of phosphorylated CREB after forskolin stimulation was much lower than in wild-type neurons. Therefore, MAP2 is an anchoring protein of PKA in dendrites, whose loss leads to reduced amount of dendritic and total PKA and reduced activation of CREB.

Introduction

Microtubule-associated protein (MAP)* 2, the most abundant MAP in the brain, mainly localizes to the somatodendritic compartments in neurons. MAP2 has three or four microtubule binding domains at the COOH-terminal, and is involved in microtubule assembly and stabilization in dendrites (Goedert et al., 1991; Matus, 1994). It is also a major component of cross-bridges between microtubules (MTs) or between MTs and other cytoskeletal components in dendrites as revealed by quick freeze deep etch microscopy and immunoelectron microscopy (Shiomura and Hirokawa, 1987; Hirokawa et al., 1988). Overexpression of MAP2 resulted in intracellular bundling of MTs and process formation of nonneuronal cells (Lewis et al., 1989; Chen et al., 1992; Takemura et al., 1992; Hirokawa, 1994). Inhibition of MAP2 production by the addition of MAP2 antisense oligonucleotides into the culture medium of cerebellar macroneurons (Caceres et al., 1992) or the expression of antisense construct in the embryonic carcinoma cell lines resulted in the inhibition of the initial neurite formation and outgrowth (Dinsmore and Solomon, 1991). These results indicated MAP2 is essential for dendritic growth by selectively stabilizing dendritic MTs.

MAP2 is also known to have a domain that binds to the regulatory subunit II of cAMP-dependent protein kinase (PKA) near its NH₂ terminus (Vallee et al., 1981; Theurkauf and Vallee, 1982). The catalytic subunit of PKA binds to the regulatory subunit I (RI) or II (RII) in an inactive state, and the inactive complex exists as a heterotetramer (two catalytic and two regulatory subunits). The catalytic subunits dissociate from RI or RII when intracellular cAMP binds to the cAMP binding sites of RI or RII. In neurons, RII is the major regulatory subunit and comprises 85–90% of total regulatory subunit of PKA (Stein et al., 1987; Ventra et al., 1996). Thus, the localization of RII determines the localization of PKA itself in neurons, whose kinase activity is known to be involved in various biological functions in neurons, including the formation of long-term memory in mice (Abel et al., 1997; Malleret et al., 2001). As MAP2 is highly enriched in the dendritic cytoskeleton, it has been assumed that it is one of the major PKA-anchoring proteins in dendrites, and thus assumed to play roles in PKA signal transduction. However, the role of MAP2 in PKA signal transduction has not been intensively examined in spite of its abundance in neurons. Therefore, to know the role of MAP2 in the intracellular structure and the signal transduction by PKA in neurons, we disrupted the map2 gene by homologous recombination in embryonic stem cells and made MAP2-
deficient mice. In our previous paper (Teng et al., 2001), we analyzed MAP2 MAP1B double knockout mice and showed that the double knockout mice had decreased microtubule spacing in dendrites, decreased dendritic elongation of immature hippocampal neurons, fiber tract malformations, and defects in neuronal migration. These results indicated that MAP2 and MAP1B act synergistically in neuronal migration and dendritic extension. However, the unique role of MAP2 on MT organization and PKA signal transduction was not reported in the previous paper. In this paper, we report that MAP2 itself is important as a structural protein and as an anchoring protein of PKA in dendrites whose loss leads to reduced dendritic and total PKA and reduced activation of cAMP-responsive element binding protein (CREB).

Results

Targeting the map2 gene
To disrupt the endogenous map2 gene, a targeting vector was constructed (Harada et al., 1998) with the neo gene inserted into the fifteenth exon (Kalcheva et al., 1995) that encodes the first MT binding domain and the preceding proline-rich domain (Teng et al., 2001). Homozygous mutant (map2−/−) animals were fertile and have shown no evidence of premature mortality, but they were smaller and 10–20% less in body weight than their littermate controls. However, their brain weight was not significantly less than their littermate controls.

Microtubule density is reduced and fine structures are altered in dendrites of MAP2-deficient mice
To further investigate the morphological changes in map2−/− brains, we examined cerebellar tissue by electron microscopy. The gross cytoarchitecture and cell morphology of map2−/− cerebellum were indistinguishable from those of the control cerebellum. Our previous paper (Teng et al., 2001), reported decreased spacing between MTs in neurites of MAP2 MAP1B double knockout, but it did not report quantitation of the MT density of map2−/− dendrites. When we examined Purkinje cell dendrites, MT density in map2−/− mice (35.6 ± 8.5 MTs/μm² [mean ± SD]; n = 31) is 23% less than that in wild-type mice (46.3 ± 8.6; n = 25) (P < 0.001, Student’s t test). Therefore, though MAP2 and MAP1B act synergistically in MT organization in dendrites as reported previously, the polymerization of dendritic MTs is affected by the absence of MAP2 alone.

Also in our previous paper (Teng et al., 2001), we were not able to detect changes in MT organization in map2−/− dendrites using conventional electron microscopy. Therefore, to further assess the structural changes in the dendrites of map2−/− mice, cerebellar Purkinje cell dendrites of map2−/− and wild-type mice were examined by quick-freeze, deep-etch electron microscopy, which was more suitable to examine cytoskeletal organization (Harada et al., 1994). In wild-type dendrites, we observed thick filamentous cross-bridges between MTs (Fig. 1, arrows), confirming our previous studies (Hirokawa et al., 1988; Chen et al., 1992). In map2−/− dendrites, the cross-bridges between MTs (Fig. 1, arrows) were thinner (for wild type, 7.08 ± 0.36 nm, 45 cross-bridges from two animals; for knockout, 4.38 ± 0.63, 28 cross-bridges from two animals) (P < 0.01). Globular structures were observed on the surface of MTs in wild-type dendrites (Fig. 1, arrowheads), but these globular structures were rarely observed in mutant dendrites. These observations led us to conclude that MAP2 was required to form thick cross-bridges between MTs, and that MAP2 was required for MT organization in the dendrites.

Dendrite length is reduced in MAP2-deficient neurons in vitro and in vivo
To examine the effect of MAP2 in neuronal maturation, we observed hippocampal neurons in culture. In our previous paper, we did not detect significant difference in the polarization process at days 0.5, 1, 1.5, 2, and 3 after plating between map2−/− and the control neurons (Teng et al., 2001). However, we did not report changes in map2−/− hippocampal culture at later stages. In our previous paper (Teng et al., 2001), the length of neurites of long-term cultured neurons could not be measured because we plated neurons so densely that we were unable to discriminate each neurite. Therefore, we plated the hippocampal neurons at a lower density and cultured for a long period (3 wk), the length of map2−/− dendrites was significantly (~30%) shorter than that of the control dendrites (Fig. 2, A and C): for WT1, 153 ± 54 μm n = 25 neurons; for WT2, 135 ± 50 μm n = 34; for WT3, 144 ± 58, n = 30; for KO1, 98 ± 34, n = 22; for KO2, 106 ± 35, n = 35; and for KO3, 92 ± 27, n = 33 (P < 0.00001, Post-hoc test). These results indicate that MAP2 is required for the dendrite extension at later stages in culture.

To assess the morphology of the dendritic trees in map2−/− hippocampal neurons in vivo, we stained hippocampal tissue by using the rapid Golgi method. When we observed the CA1 region of the hippocampus, apical dendrites of map2−/− hip-
Hippocampal neurons were shortened compared with wild-type dendrites, whereas the basal dendrites were not affected (Fig. 2 B). The length of mutant apical dendrites was 22% shorter than that of wild-type dendrites (Fig. 2 D), whereas the length of basal dendrites did not differ (for map2−/− mice, 122 ± 10 μm; 11 dendrites from 11 tissue sections [we selected the longest dendrite from one section] from two animals; for wild-type mice, 122 ± 11; the number of samples and animals are the same as those of knockout) (Fig. 2 D). Therefore, map2 is required for the morphology of the dendrites both in vivo and in vitro.

**Various PKA subunits are reduced in map2−/− dendrites of hippocampal and cortical neurons**

Next, we examined the localization of various PKA subunits in hippocampal neurons from map2−/− mouse. (A) Hippocampal cells in culture for 3 wk stained with rhodamine phalloidin. Dendrites of map2−/− cells (KO) are shorter than wild-type cells (WT). Bar, 100 μm. (B) Pyramidal neurons in CA1 area of hippocampal tissues stained by the rapid Golgi method. Apical dendrites of map2−/− neurons (KO) are shorter than those of wild-type neurons (WT). However, the length of basal dendrites of map2−/− neurons is similar to that of wild-type neurons. Bar, 100 μm. (C) Quantitation of the average length of dendrites between wild-type (WT1–3) and map2−/− cells (KO1–3). The average lengths of map2−/− dendrites contained within a neuron are significantly shorter than those of wild-type dendrites (mean ± SD). P < 0.00001 (Post-hoc test). (D) Quantitation of the average length of apical and basal dendrites between wild-type (WT) and map2−/− cells (KO). The average lengths of map2−/− apical dendrites contained within a neuron are significantly shorter than those of wild-type dendrites (for map2−/− mice, 179 ± 15 μm; 11 dendrites from 11 tissue sections from two animals [we selected the longest dendrite from one section]; for wild-type mice, 229 ± 22; the number of samples and animals are the same as those of knockout) (mean ± SD). P < 0.001 (Post-hoc test).

Levels of PKA (RII α, β, and the catalytic subunits) The dendritic PKA staining is more apparent in early postnatal ages (2 wk of age; Fig. 3, A, A′, C, C′, E, E′, G, and G′), but is also observed in adult mice (8 wk of age; Fig. 4, A, B, and D). In addition to this dendritic staining, the mossy fiber axons (Fig. 3, mf) from the granule cells of dentate gyrus (Fig. 3, stained by anti-τ antibody in E′′, F′′, G′′′, and H′′′) contained RII β and the catalytic subunits (Fig. 3, A, A′, C, C′, G, and G′ and Fig. 4 A). RII α staining was very low in the mossy fiber axons (Fig. 3, E and E′). However, in map2−/− hippocampus, the dendritic staining of RII α, β, and the catalytic subunits reduced significantly (Fig. 3, B–B′′, D–D′′, F, F′, H, and H′ and Fig. 4, A′, B′, D′), whereas the axonal staining seemed relatively unaffected as revealed by the staining of mossy fiber in Fig. 3 (B and B′, mf). The axonal staining also seemed unaffected in the fimbria that includes axons from the hippocampal pyramidal neurons (unpublished data).

To confirm the reduction of various PKA subunits in map2−/− dendrites, we stained cerebral cortex. In the control cerebral cortex of adult mice, strong signals for PKA subunits were observed in neurons throughout the cortex, but it accumulated primarily in the apical dendrites and the neurons.
soma in the pyramidal neurons (Fig. 5, A–C). In map2−/− cortex, the signals in the soma were relatively unchanged, but those in the dendrites were significantly reduced (Fig. 5, A′–C′). Similar to the observation in the hippocampus, the content of RII β and the catalytic subunits seemed unchanged in the axonal compartments that were stained by tau-1 antibody (unpublished data).

To examine the content and the subcellular localization of various PKA subunits, we stained dissociated hippocampal cultured neurons. In hippocampal neurons, the staining of

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**Figure 3.** Reduced expression of PKA in the dendrites of 2-wk-old wild-type and map2−/− hippocampus. Low (A–H) and high (A′–H′) magnification view of CA3 region stained with anti–RII β monoclonal antibody (A, A′, B, and B′), anti–catalytic subunit antisera (C, C′, D, and D′), anti–RII α antisera (E, E′, F, and F′), and anti–RII β antisera (G, G′, H, and H′). High magnification view of CA1 region stained with anti–RII β monoclonal antibody (A″ and B″) and anti–catalytic subunit antisera (C″ and D″). E″, F″, G″, and H″ are the same sections as E′, F′, G′, and H′, respectively, stained with anti–τ antibody (tau-1) to show the localization of mossy fiber (mf). Arrows in A′, A″, C′, and C″ indicate the dendritic staining of PKA in wild-type pyramidal cells. Bars, 100 μm.

**Figure 4.** Reduced expression of various subunits of PKA in the dendrites of adult wild-type and map2−/− hippocampus. CA3 region (A and A′) and CA1 region (B–D and B′–D′) stained with anti–RII β monoclonal antibody (A and A′) and anti–catalytic subunit antisera (B and B′). The same sections stained with anti–catalytic subunit antisera (B and B′) were stained with anti–MAP1A monoclonal antibody (C and C′) to show the localization of dendrites and the merged figures between PKA catalytic subunit (green) and MAP1A (red) are shown in D and D′. Catalytic subunit is reduced in the dendrites of map2−/− hippocampal neurons. Bar, 100 μm.
PKA subunits (RIIα, β and the catalytic subunit) was weaker in the somatodendritic compartments of map2−/− neurons (Fig. 6, A′–E′) than wild-type neurons (Fig. 6, A–E), whereas the staining of Ca/calmodulin-dependent kinase in map2−/− neurons (Fig. 6 F′) was similar to that in wild-type neurons (Fig. 6 F). The staining of PKA appeared linear and concentrated in the center of the dendrites in the control neurons at higher magnification (Fig. 6 B). In contrast, the staining of PKA in map2−/− appeared punctate, and thus did not appear to be associated with cytoskeletal elements. Since PKA is known to bind to MAP2, it was assumed that PKA was colocalized with MTs in the dendrites. To know whether PKA was associated with cytoskeletal compartments, we treated the cultured neurons with 0.1% Triton X-100. In the control neurons, the amount and the distribution of PKA were similar to the staining without detergent extraction (Fig. 6 C). However, the punctate PKA staining in the dendrites and the staining in the soma were greatly reduced in map2−/− neurons (Fig. 6 C′), suggesting the remaining PKA molecules in the somatodendritic compartments in map2−/− neurons were not associated with cytoskeletal compartments. We also observed hippocampal and cortical neurons well separated from the neighboring neurons to examine the amount of PKA in the axonal or dendritic compartments. The staining in map2−/− dendrites was clearly reduced, but axonal staining was relatively unchanged (unpublished data). In addition, to know the localization and the amount of PKA more precisely at the subcellular level, immunoelectron microscopy was performed in cultured hippocampal neurons. When wild-type neurons were stained with anti-RII β monoclonal antibody and the secondary antibody conjugated with gold particles, followed by silver enhancement, many electron-dense particles were associated with MTs in the dendrite. However, in map2−/− neurons, the number of signals reduced significantly. When we quantitated these signals, the density of signals in map2−/−
Figure 7. Reduced amount of PKA and reduced activation (phosphorylation) of CREB after forskolin induction in map2−/− mice. (A) Western blots of the microtubule pellet fraction (MT ppt), the crude extracts from the hippocampal tissue (HIP tissue), and the crude extracts from the cultured neurons (HIP culture) probed with anti-RII β monoclonal antibody (RIIβ) and anti-catalytic subunit antiserum (Cat). (B) Quantitation of PKA (RIIβ and Cat) in MT ppt (n = 3; one sample was taken from one animal), HIP tissue (n = 3), and HIP culture (n = 3). The relative amounts of map2−/− against wild-type samples are shown. (C) Western blots of phosphorylated CREB. Wild-type (WT) and map2−/− (KO) neurons were homogenized before (F−) and 30 min after forskolin stimulation (F+), and equal amounts of samples from wild-type and knockout mice were loaded and separated with polyacrylamide gel. (D) Quantitation of the results of immunoblot in C. The amount of phosphorylated CREB is increased about threefold in wild-type neurons (WT), whereas the amount is nearly unchanged in map2−/− neurons (KO). The number of samples and animals are the same as in B.

Total amount of PKA is also reduced in map2−/− hippocampal tissue and cultured neurons, and the rate of increase in phosphorylated CREB is less than wild-type neurons after forskolin stimulation

As the amount of PKA was decreased in the somatodendritic compartments of map2−/− neurons in vitro and in vivo from the observation of immunofluorescence microscopy and immunoEM, we performed quantitative immunoblot of various fractions with antibodies that recognized RII β and the catalytic subunits of PKA (Fig. 7, A and B). Quite surprisingly, the contents of PKA were greatly reduced in the MT pellet fraction (RII β, ~5%; catalytic subunit, ~7% of wild type). Similar reduction was observed when the activity of PKA was quantitated (unpublished data). In addition, PKA was also reduced in the crude homogenate of hippocampal tissues (RII β, ~70%; catalytic, ~80% of wild type). When we used the crude extracts from the cultured hippocampal neurons, the PKA content was much more reduced (RII β, ~50%; catalytic subunit, ~60% of wild type). These results indicate that the total amount of PKA was reduced in map2−/− neurons probably because the PKA molecules anchored and pooled in the MT-rich regions in the dendrites were significantly reduced. To assess possible effects from the reduction of PKA in map2−/− neurons, we measured the extent of phosphorylation of a PKA substrate. We quantitated the amount of phosphorylation at Ser133 of CREB, a transcription factor activated by PKA, in cultured hippocampal neurons before and after PKA stimulation by forskolin (Fig. 7, C and D). Before stimulation, the amount of phosphorylated CREB at Ser133 in map2−/− neurons was about twofold of that in wild-type neurons. However, the amount of phosphorylated CREB was relatively unchanged (~1.1-fold increase) even after forskolin stimulation. In contrast, the amount of phosphorylated CREB increased about threefold by stimulation in wild-type neurons. The amount of total CREB did not differ significantly between map2−/− and wild-type neurons (unpublished data). From these results, MAP2 is required for controlling the total amount of PKA and proper induction of activated (phosphorylated) CREB.

Discussion

The role of MAP2 in dendrite elongation

MAP2 promotes MT polymerization in vitro. When overexpressed in nonneuronal cells (Lewis et al., 1989; Chen et al., 1992), MAP2 promotes the intracellular MT bundling, and when introduced into Sf 9 insect cells by baculovirus infection, it promotes the formation of MT bundles and the neurite-like process formation (Chen et al., 1992). MAP2 was also considered to be essential in the formation of the dendrites because MAP2 antisense exposure in neuronal cell culture specifically inhibits the elaboration of dendrite-like processes (Caceres et al., 1992).

We observed a reduction in MT density in dendrites of MAP2-deficient mice, which was not described in our previous paper (Teng et al., 2001). Although we did not measure MT density in the dendrites of MAP2 and MAP1B double knockout mice, MAP2 alone is effective in MT polymerization in dendrites from this result. We also observed reduced...
dendritic outgrowth. We observed a great reduction of PKA in the MT pellet fraction of CREB after forskolin stimulation leads to the reduction in the total amount of CREB. The possible explanation for this contradiction is as follows: in vivo, various external factors (growth factors, interaction with other neurons, etc.) may enhance the growth of dendrites and partially compensate for the reduced dendritic extension by the loss of MAP2 in map2−/− neurons. However, in dissociated culture where these external factors do not exist, map2−/− neurons may show their reduced internal activity in dendritic outgrowth.

**Significant reduction of PKA in MT-rich region leads to the reduction in the total amount of PKA in neurons and reduced activation of CREB after forskolin stimulation**

We observed a great reduction of PKA in the MT pellet fraction of map2−/− neurons (~1/20 [RII β] and 1/15 [catalytic subunit] of wild type) by quantitative immunoblotting. This was confirmed both by measuring activities of PKA in the MT pellet and by immunoelectron microscopy, where the PKA (RII) signals in the MT-rich region of map2−/− dendrites were not significantly different from nonimmune controls. As a result of this reduction, the total amounts of PKA in map2−/− hippocampal neurons were decreased to about half of those of wild-type neurons. There is a tendency that the amount of the catalytic subunit is less affected than that of RII. This is probably due to the presence of the RI that do not bind to MAP2. Although the amount of RI is much less than RII in neurons and astrocytes, the former is nearly equal to the latter in oligodendrocytes (Stein et al., 1987). This also explains why the reduction rate of PKA in hippocampal tissue is less than that in dissociated hippocampal neurons. In MAP2-deficient neurons, the outer membranous (or submembranous) structures such as the postsynaptic spines and the growth cones, and the internal membranous structures such as the Golgi complex, seem to be the main region of PKA localization from previous papers (De Camilli et al., 1986) and from our immunofluorescence microscopy. In this paper, the localization of PKA seemed to shift from the MT-rich regions to the soluble fraction or these membranous structures. However, considering that the total amount of PKA was still reduced in map2−/− neurons, the regions that bound to PKA in map2−/− neurons were not sufficient for anchoring all the PKA molecules released from the MT-rich region in the absence of MAP2. This observation strongly indicates that MAP2 is the main PKA-anchoring protein in dendrites. However, in the axonal compartments, the amount of PKA (RII β and the catalytic subunits) did not change significantly as in the dendritic compartments from the observation by immunofluorescence microscopy. This implies that other molecules anchor PKA, especially RII β, in the axons. This may relate to the fact that the affinity of RII β to MAP2 is less than that of RII α. Therefore, RII α seems to be designed to anchor the PKA catalytic subunits to MAP2 in the dendrites, whereas RII β seems to be designed to anchor them to MAP2 in the dendrites and to other anchoring proteins in the axons.

We also observed an increase in the basal phosphorylation of CREB in map2−/− neurons compared with wild-type neurons. Phosphorylated CREB only slightly increased after forskolin stimulation in map2−/− neurons. Thus, the role of MAP2 seems to trap and store PKA in the dendritic MTs before forskolin stimulation. This role enables PKA to be rapidly released into the cytoplasm to phosphorylate a number of substrates after stimulation.

PKA is also known to phosphorylate a number of substrates including cytoskeletons (Hisanaga et al., 1994) and microtubule-dependent motor proteins (Sato-Yoshitake et al., 1992). A MAP2 knockout mouse can be used for investigating the role of PKA in dendrites by analyzing the phosphorylation of these substrates.

**Materials and methods**

**Hippocampal cell culture**

Methods for preparing the hippocampal cell cultures followed those described previously (Harada et al., 1994). Usually, in one set of experiments, fetuses belonging to the same litter were used, the same conditioned medium was used, and feeder glial cells were dissociated from littermate embryos. We also used hippocampal neurons cultured without glial cells for immunofluorescence, immunoelectron microscopy, and for Western blot. The result obtained with either culture protocol was similar therefore, the results were pooled. We followed the neurons photographically and measured the length and counted the number of dendrites as described previously (Harada et al., 1994; Teng et al., 2001). To compare the length of dendrites, we first calculated the average length of dendrites of a neuron by averaging the lengths of all visible dendrites and then obtained the average in each animal by using the average of each neuron.

**Immunoblotting**

For preparation of crude extracts, the hippocampus was dissected and homogenized with 25 mM Tris, pH 6.8, and 2% SDS, followed by sonication and boiling to lower the viscosity. Cultured cells were lysed in the same buffer and treated in the same manner. Crude extracts were made by centrifuging the homogenates at 20,000 g for 15 min at 4°C. Preparation of MT pellet fraction was performed as described previously (Sato-Yoshitake et al., 1989). In brief, the brains were dissected out and homogenized in 3 vol PEM (100 mM PIPES, 1 mM EDTA, and 1 mM MgCl2) including 1 μM PMSF. Crude extracts were prepared by centrifuging the homogenates at 100,000 g for 15 min at 4°C. A microtubule polymerizing agent, taxol, was added to the crude extracts to a final concentration of 20 μM. The crude extracts were incubated at 37°C for 15 min, centrifuged at 100,000 g for 30 min at 27°C and the resultant pellet was named MT ppt. Protein concentration was determined by BCA protein assay reagent (Pierce Chemical Co.). Equal amounts of crude extracts and MT ppt were loaded and separated with polyacrylamide gel. Proteins were electrophoretically transferred to nylon filters (Millipore). Nylon filters with transferred brain proteins were blocked with 5% skim milk in TBS, incubated in monoclonal antibodies for 1 h, rinsed in TBS and TBS containing 0.05% Tween 20, followed by incubation in rabbit anti–mouse IgG, and incubated for 1 h with 125I-labeled protein A (Amersham Biosciences). Binding was detected by autoradiography using an imaging analyzer (model BAS-2000; Fuji Film). For detection of phospho-CREB, alkaline phosphatase–conjugated secondary antibody was used because the signals obtained by 125I-labeled protein A were not strong enough for quantitation. In this case, quantitation was performed by scanning the Western blot of various amounts of map2−/− and control samples. Measurement of the intensity of the bands was performed by NIH
Immunocytochemical analysis

2-, 4-, and 8-wk-old mice were anesthetized with ether and nembutal (WAKO) and perfused with 2% paraformaldehyde in 0.1 M phosphate buffer, pH 7.4. Various parts of brains (cerebellum, cerebral cortex, striatum, hippocampus, and spinal cord) were dissected and stored in the same fixative overnight at RT. The procedure for cryoprotection, freezing, and sectioning was described previously (Harada et al., 1990). Sections were allowed to react with the primary antibodies described previously (Harada et al., 1990) followed by the secondary antibodies (Alexa 488 and 568-conjugated goat anti-mouse or goat anti-rabbit IgG and Alexa 633-conjugated phalloidin (Molecular Probes). In case of cultured hippocampal neurons, neurons on coverslips were fixed with 2% paraformaldehyde in PBS, pH 6.8, permeabilized in 0.1% Triton in PBS for 5 min at RT, and incubated in the primary and the secondary antibodies as described above. Samples were observed using a laser scanning confocal microscope (model LSM510; ZEISS).

Rapid Golgi staining

Mice were perfused with 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.4. Hippocampus was dissected and further fixed for 2 h at RT. The dissected hippocampus was washed twice in 3% potassium dichromate in water and incubated in 5% glutaraldehyde and 2.5% potassium dichromate in water at 20°C for 4–5 d. Tissues were washed several times in 0.75% silver nitrate in water and then incubated in 0.75% silver nitrate in water at 20°C for 4–5 d again. 100–150-μm thick slices were obtained by cutting the tissue perpendicular to the long axis of hippocampus with a microtome (model DTK-1000; Dosaka EM). The Golgi-stained neurons were observed with a laser scanning confocal microscope by the reflection contrast mode. 10 optical sections (distances between slices were 10 μm) were obtained from one hippocampal slice, and the images were merged to obtain an image as complete as possible of each dendritic tree. To calculate the length of dendrites, we selected three dendrites from one section that appeared to fully extend from the cell body to the tip of the dendrite and use the longest dendrite to represent the section. We took an average of the length of representatives within each genotype (wild type or knockout).

Conventional electron microscopy

Mice were perfused with 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.4. Tissues (hippocampus, cerebellum, and olfactory bulb) were dissected and further fixed for 2 h at RT. Tissues (hippocampus, cerebellum, and olfactory bulb) were dissected and further fixed for 2 h at RT. Tissues were washed several times in 0.75% silver nitrate in water and then incubated in 5% glutaraldehyde and 2.5% potassium dichromate in water at 20°C. Tissues were then washed and postfixed in 4% glutaraldehyde and 1% osmium tetroxide in 0.1 M cacodylate buffer, pH 7.4. Tissues were then dehydrated through an acetone series and embedded in Epon. Thin sections were obtained with a diamond knife on a Reichert Ultracut EM and stained with 2% uranyl acetate in 50% ethanol and 0.5% lead citrate. Sections were observed using a transmission electron microscope (model Hitachi H7600). Samples were observed using a scanning transmission electron microscope (model JSM-6490LV; JEOL).

Quick-freeze, deep-etch electron microscopy

Cerebellar tissues were dissected, and thin sagittal slices of tissue were saponin-treated and processed for quick-freeze, deep-etch electron microscopy as described previously (Hirokawa et al., 1988). For quantification of cross-bridges, two knockout and two wild-type mice were used. Since the length and diameter of cross-bridges between MTs were similar within each genotype, the values were pooled.

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