Caldesmon Regulates Axon Extension through Interaction with Myosin II

Tsuyoshi Morita¹, Taira Mayanagi¹,² and Kenji Sobue¹,²

¹From the Department of Neuroscience, Osaka University Graduate School of Medicine, 2-2 Yamadaoka, Suita, Osaka 565-0871, Japan
²Department of Neuroscience, Institute for Biomedical Sciences, Iwate Medical University, 2-1-1 Nishitokuta, Yahaba-cho, Shiwa-gun, Iwate 028-3694, Japan

Running title: Caldesmon regulates axon extension

To whom correspondence should be addressed: Kenji Sobue, Department of Neuroscience, Institute for Biomedical Sciences, Iwate Medical University, 2-1-1 Nishitokuta, Yahaba-cho, Shiwa-gun, Iwate 028-3694, Japan, Tel.: +81-19-651-5710; Fax: +81-19-908-8020; E-mail: ksobue@iwate-med.ac.jp

keywords: caldesmon; axon extension; myosin

Background: Axon extension, an essential step for creating neural circuits, is regulated by cytoskeletal dynamics.

Results: Caldesmon is a regulator of the actin cytoskeleton and enhances axon extension through direct interaction with myosin II.

Conclusion: Caldesmon binding to myosin II inhibits myosin II function, resulting in the enhancement of axon extension.

Significance: This study elucidates how caldesmon-regulated actin-myosin system is involved in axon extension.

SUMMARY

To begin the process of forming neural circuits, new neurons first establish their polarity and extend their axon. Axon extension is guided and regulated by highly coordinated cytoskeletal dynamics. Here we demonstrate that in hippocampal neurons, the actin-binding protein caldesmon accumulates in distal axons, and its N-terminal interaction with myosin II enhances axon extension. In cortical neural progenitor cells, caldesmon knockdown suppresses axon extension and neuronal polarity. These results indicate that caldesmon is an important regulator of axon development.

Neurons in the developing brain extend axonal and dendritic arbors that create a complex circuitry, and the guided extension of axonal fibers is an essential step in this process. Axon extension is regulated by the coordinated interaction of microtubules and actin filaments.
in the axonal growth cone. A growing body of evidence indicates that microtubule polymerization and stabilization play positive roles in axon extension (1), whereas actin filament roles are more complicated. For example, knocking out Ena/VASP or Cdc42, which positively regulate actin polymerization, causes axonal tract loss (2, 3). In contrast, inhibiting the actin nucleation factor Arp2/3 and pharmacologically destabilizing actin filaments enhances axon extension (4, 5). Thus, the fundamental details of axon guidance and regulation by actin filaments are not well understood.

Caldesmon (CaD) was first identified as a smooth-muscle protein that binds calmodulin and actin (6). It has since been found to be ubiquitously expressed in smooth muscle and non-muscle cells, and to regulate Ca$^{2+}$-dependent actomyosin contraction (7, 8). CaD binds to the side of filamentous actin (F-actin) and inhibits actin-myosin interactions, as revealed by superprecipitation assays and actin-activated myosin ATPase activity (9-11). CaD binding also stabilizes F-actin filaments by enhancing actin–tropomyosin binding and preventing the actin-severing activity of gelsolin or coflin (12, 13). CaD plays important roles in migration of non-muscle cells via regulating actin-myosin system (8). We recently reported that CaD is involved in detrimental glucocorticoid-induced effects during cortical brain development (14, 15): glucocorticoids increase CaD levels, transiently retarding the radial migration of cortical neuronal progenitor cells. We also reported that CaD localizes to neuronal growth cones (16). Thus, it seems that CaD plays multiple important roles in neuronal development. In this report, we demonstrate a novel role for CaD in axon extension via its N-terminal myosin-binding sequence.

**EXPERIMENTAL PROCEDURES**

**Materials** – The myosin II ATPase inhibitor blebbistatin, the myosin light chain kinase inhibitor ML-7, and the Rho-associated protein kinase inhibitor Y27632 were purchased from Merck. The following antibodies were purchased: anti-tau1 (Chemicon), anti-MAP2 (Chemicon), anti-nonmuscle myosin IIA (Abcam), anti-nonmuscle myosin IIB (Abcam), anti-GFP (Invitrogen), anti-FLAG (Sigma), anti-myc (9E10, Santa Cruz), and anti-GAPDH (FL-335, Santa Cruz). Anti-CaD antibody was generated as previously described (17).

**Cell culture and immunostaining** – Hippocampal neurons were prepared from rat hippocampi on embryonic day 18.5. The dissociated neurons were plated on poly-L-lysine-coated coverslips, and cultured in glial-conditioned MEM containing 1 mM pyruvate, 0.6% (W/V) D-Glucose, and 2% B27 supplement (Invitrogen). The next day, the culture was changed to a neurobasal medium containing 2% B27 supplement and 0.5 mM L-glutamine. Cortical NPCs were prepared from rat cerebral cortex on embryonic day 15.5 (E15.5), cultured as previously described (14), plated on laminin-coated coverslips, and
cultured under basic FGF-free conditions to induce their differentiation into polarized neurons. A549 and HEK293T cells were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum. Cells cultured on coverslips were fixed using 4% paraformaldehyde and then processed for immunocytochemistry. To label F-actin, Alexa 568-phalloidin (Molecular Probes) was added to the secondary antibody solution.

**Transfection** – Hippocampal neurons prepared from rat embryos on E18.5 were transfected by the calcium phosphate method as described previously (18). In brief, DNA-calcium phosphate precipitates were prepared using a calcium phosphate transfection kit (Invitrogen). The hippocampal neurons were plated on a NunclonΔ surface plate (Nalge Nunc International) and incubated with the precipitates for 3 hours. The transfected neurons were replated on poly-L-lysine-coated coverslips and cultured for 3 to 5 days. A549 and HEK293T cells were transfected using Lipofectamine 2000 or Lipofectamine LTX (Invitrogen).

**Expression plasmids** – The coding regions for human l-CaD, its N-terminus (1-263 amino acids), C-terminus (264-558 amino acids), and N-terminus Δ 21-47 (lacking amino acids 21-47 from the N-terminus), and the N-terminal fragments of rat myosin IIA (1-1961 amino acids) and IIB (1-1976 amino acids) were amplified by PCR and subcloned into the highly efficient mammalian expression plasmid pCAGGS. EGFP and myc tag sequences were fused to the 5’ end of the coding sequences. The mcherry-LifeAct expression vector was constructed as previously reported (19).

**RNA interference** – Short-interfering RNAs (siRNAs) against rat CaD were transfected into growing cortical NPCs using Lipofectamine RNAi MAX (Invitrogen). MicroRNA (miRNA) plasmids against rat CaD were constructed as previously described (14) and transfected into hippocampal neurons by calcium phosphate precipitation. The targeting sequences and the siRNA and miRNA knockdown efficacy were reported in our previous studies (14, 20).

**Immunoprecipitation** – HEK293T cells with transfected expression vectors were lysed with Triton-X-buffer [0.05% Triton X-100 (pH 7.6), 30 mM Tris-HCl, 50 mM NaCl 5mM EGTA, 5 mM MgCl₂, 1 mM ATP, and protease inhibitor cocktail for use with mammalian cell and tissue extracts (Nacalai Tesque)]. Immunoprecipitation was performed using the earlier-listed antibodies and protein G-Sepharose (GE Healthcare Life Sciences). The Sepharose beads were boiled in SDS-sample buffer to elute the immunocomplexes.

**RESULTS**

*CaD enhances axon extension in hippocampal neurons* – CaD, a ubiquitous regulator of the actin cytoskeleton, localizes along actin fibers and in the ruffling membrane...
Caldesmon regulates axon extension

(7, 8). Here, we found that CaD was located in the soma and growth cones of primary cultured hippocampal neurons, with the strongest expression in the distal axon (Fig 1A-D). CaD levels increased for 3 to 7 days \textit{in vitro} (DIV) (2.3 ± 0.8-fold at 7 DIV vs. 2 DIV) while the neurons established polarity and actively extended axons (Fig 1E). The location and time-course of CaD’s expression in these cells are consistent with its having a role in axon extension.

We therefore investigated CaD’s function in neurite outgrowth by overexpressing or knocking down CaD in hippocampal neurons. We used GFP-fused CaD (GFP-CaD), which has the same functions as endogenous CaD (14, 20). GFP-CaD dramatically enhanced axon extension but did not significantly affect dendrite length as compared with the control, GFP (Fig 1F, G). GFP-CaD also enhanced formation of filopodia-like protrusions from the soma and axon branches (Fig 1F). These CaD-induced protrusions were composed of concentrated actin filaments and were distinct from the main axonal branches, which were filled with microtubules (Fig 1H). Knocking down the endogenous CaD decreased axon length, but not dendritic length (Fig 1I, J), indicating that CaD accumulates in the distal axon of hippocampal neurons during their development and enhances axon extension.

\textit{CaD regulates axon development in cortical NPCs} – To monitor CaD’s involvement in early events in neurite outgrowth, we used cortical neural progenitor cells (NPCs), which proliferate as non-polarized cells in the presence of basic fibroblast growth factor (FGF) (14, 21). Under basic FGF-free conditions, however, NPCs stop proliferating and establish neuronal polarity with MAP2-positive dendrites and a tau1-positive axon (Fig 2A). When CaD was knocked down with siRNAs in proliferating NPCs, tau1-staining showed that the establishment of neuronal polarity was significantly suppressed within three culture days under basic FGF-free conditions (Fig 2A, B). Even in polarized cells, the length of tau1-positive axons was significantly shortened by CaD knockdown (Fig 2A, C), as observed in hippocampal neurons. At an early stage of NPCs differentiation into polarized cells, immature axons were often stained with both anti-MAP2 and anti-tau1 antibodies. In the CaD-knockdown NPCs, some short axons were MAP2/tau1 double positive, suggesting delayed development of these cells. These findings indicate that CaD plays important roles in establishing neuronal polarity and in axon extension in developing NPCs.

\textit{CaD-myosin interaction required for axon extension} – CaD has been reported to bind smooth muscle myosin at its N-terminus and F-actin at its C-terminus, suggesting that it functions to link these molecules (22). In the growth cone of hippocampal neuronal axons, CaD colocalized with F-actin and myosin IIA/IIB, the major non-muscle isoforms of myosin II (Fig 3A). To examine myosin and
actin involvement in CaD-induced axon extension, CaD N- and C-terminal fragments (N-CaD and C-CaD) were expressed separately in hippocampal neurons. N-CaD enhanced axon extension like full-length CaD, but C-CaD did not (Fig 3B-D), suggesting that CaD’s interaction with myosin, but not F-actin, is necessary for CaD-induced axon extension. On the other hand, C-CaD, but not N-CaD, induced formation of the filopodia-like protrusions like full-length CaD (Fig. 1F and Fig. 3C, E), suggesting that this effect is dependent on the C-terminal actin binding domains.

Co-immunoprecipitation was used to determine whether non-muscle myosin II, like smooth- and skeletal-muscle myosins, binds to CaD. Because CaD is reported to bind to the S-1 and S-2 regions of smooth and skeletal muscle myosins (23), we examined CaD interactions with myosin IIA or IIB N-terminal fragments, which are composed of a globular head domain, a neck region, and a small tail fragment corresponding to heavy meromyosin (HMM). As with smooth- and skeletal-muscle myosins, CaD bound to HMM IIA and IIB, and CaD’s C-terminal F-actin-binding domains were not necessary for these interactions (Fig 4A-C).

Previous studies demonstrated that the 27-amino acid sequence in CaD’s N-terminus (Tyr-21 to Lys-47 in human l-CaD) is necessary for binding to smooth-muscle myosin (24). N-CaD Δ21-47 fragment, in which this 27-amino acid sequence is deleted, did not interact with HMM IIA, and a CaD fragment including amino acids 1-47 was the minimum required for HMM IIA binding (Fig 4C, D). Importantly, N-CaD Δ21-47 fragment completely lost the ability to enhance axon extension (Fig 3C, D), strongly supporting the idea that CaD is accumulated in the growth cone as an actomyosin component and enhances axon extension through direct interaction with non-muscle myosin II.

**N-CaD exhibits the same effect as blebbistatin** – To determine the significance of CaD’s interaction with myosin, N-CaD or C-CaD was transfected into A549 cells. CaD has been reported to stabilize actin filaments via its C-terminal F-actin-binding domains, causing thick actin fibers to form (25, 26). In A549 cells, C-CaD strongly induced thick actin fiber formation (Fig 5A, B). On the other hand, cells expressing N-CaD showed significant actin fiber loss and a flat cell shape with prominent lamellipodia (Fig 5A, B). These effects were completely lost in A549 cells expressing an N-CaD Δ21-47 fragment lacking the 27-amino acid myosin-binding sequence (Fig 5A, B). Further, these morphological changes were very similar to those found in cells treated with the myosin II-inhibitor blebbistatin (Fig 5A). These results suggest that CaD binds to myosin at its N-terminus, and that it inhibits myosin II function independently of its C-terminal F-actin-binding domains.

**CaD changes growth cone morphology and myosin II localization** – To determine the function of CaD in growth cones, we observed
growth cone morphology and myosin II localization in the hippocampal neurons expressing CaD fragments (Fig. 6). N-CaD inhibited lamellipodia expansion, whereas C-CaD enhanced filopodia formation in growth cones. Full-length CaD induced both lamellipodia retraction and filopodia formation. N-CaD Δ21-47 had no effect on growth cone morphology.

In GFP-N-CaD-transfected neurons, myosin II staining was slightly diffuse, but distinctly strong in the basal region of the lamellipodia-poor growth cones. In the cells transfected with GFP-CaD and GFP-C-CaD, myosin II was tightly associated with filopodia. N-CaD Δ21-47 had no effect on myosin II localization. These results indicate that C-terminal actin binding domains enhances actin bundling in growth cones, leading to filopodia formation, with which myosin II associates. On the other hand, N-terminal myosin-binding domain inhibits lamellipodia formation in growth cone, but scarcely have an effect on the myosin II localization.

**CaD enhances axon extension by inhibiting myosin** – To examine how inhibiting myosin function would affect axon extension, hippocampal neurons were incubated with blebbistatin, myosin light chain kinase inhibitor ML-7, or Rho-associated protein kinase inhibitor Y27632, drugs that directly or indirectly inhibit myosin function. All of these drugs, especially the direct inhibitor blebbistatin, significantly increased axonal length compared with the vehicle control (Fig 7A, B). In GFP-CaD-transfected hippocampal neurons, however, blebbistatin did not further accelerate axon extension (Fig 7C, D). Coupled with its effects on axon extension, blebbistatin induced morphological changes in the axonal growth cones, inducing a switch from lamellipodial to filopodia-like protrusions (Fig. 7E). In the GFP-CaD-transfected neurons, axonal growth cones displayed a filopodia-like morphology without expanded lamellipodia, and their morphology was not affected by blebbistatin treatment (Fig 7E). These findings indicate that blebbistatin and CaD enhance axon extension via the same pathway, through which myosin II function is inhibited.

**DISCUSSION**

CaD is a ubiquitous regulator of the actin cytoskeleton. Most of CaD’s functional domains that bind F-actin, tropomyosin, and calmodulin are located in its C-terminus, and the C-terminal fragment can inhibit myosin ATPase activity and stabilize actin filaments (8, 25-28). CaD’s N-terminal region also has a myosin-binding sequence, through which CaD binds to smooth and skeletal muscle heavy meromyosins (23). This binding domain is probably involved in tethering myosin to actin filaments (22, 29), but the significance of myosin binding to CaD had been unclear. In our present study, we clearly demonstrated that CaD enhances axon extension through direct interaction with non-muscle myosin II via its N-terminal myosin-binding sequence. N-CaD, which lacks the all
C-terminal functional domains, exhibited the same effect on axon extension as full-length CaD (Fig 3C, D), indicating that axon extension does not depend on the CaD-mediated physical bridge between myosin and actin.

In addition to axon extension, CaD induced formation of the filopodia-like protrusions from soma and axon branches (Fig. 1F and Fig. 3E). CaD also enhanced filopodia formation in growth cones (Fig. 6). C-terminus of CaD, but not N-terminus, was required for both functions. C-terminus contains some actin binding domains, which are necessary for stabilization of actin bundles (7, 8), and the filopodia-like protrusion were composed of concentrated actin filaments (Fig. 1F). These indicate that actin stabilization by the C-terminal domains facilitates formation of these filopodial protrusions independently of N-terminal myosin binding domain.

Results of our experiments using myosin II ATPase inhibitor blebbistatin strongly suggest an inhibitory effect of N-CaD on myosin II function in hippocampal neurons and non-neuronal A549 cells (Fig 5, 7). However, previous in vitro study showed that the CaD1-597 fragment, which lacks the C-terminal actin-binding domains, does not inhibit actin-activated myosin ATPase activity via its C-terminal F-actin-binding domains, by preventing the myosin head from binding to actin in vitro (21). Considering the discrepancy between these in vitro studies and our in vivo study, we propose that N-CaD inhibits myosin II function by unknown mechanisms, which may include interacting with or recruiting additional myosin-inhibitory factors. Further investigations are required to clarify how N-CaD inhibits myosin II function.

Growing evidences indicate that myosin II function is important for axon outgrowth and axon guidance (30-35). However, the molecular mechanism underlying actomyosin-mediated axon extension has not been fully evaluated. An early study by Letourneau et al (1987) clearly demonstrated that both “push” by microtubules and “pull” by actomyosin in the growth cone play central roles in axon extension (36). Actually, actin destabilization by cytochalasin D or ADF/cofilin and myosin II inhibition by blebbistatin enhance axon extension (5, 37, and our present study). CaD may inhibit the traction force generated by the actomyosin contraction, thereby augmenting the pushing force from microtubule extension.

REFERENCES

Caldesmon regulates axon extension

Biol. 125, 359-368

FOOTNOTES
This work was supported by a Grant-in-aid for Scientific Research 20240038 and 23110510 from the Japan Society for the Promotion of Science (K. S.).
The abbreviations used are: CaD, caldesmon; DIV, day in vitro; NPCs, neural progenitor cells; FGF, fibroblast growth factor; HMM, heavy meromyosin

FIGURE REGENDS
FIGURE 1. CaD involvement in axon extension. A, CaD protein localization in primary cultured hippocampal neurons. The neurons were fixed and triple-stained with anti-CaD, anti-tau1 antibodies, and phallolidin (F-actin). Bar, 50 µm. B, Dendrite and axon fluorescence intensities measured in the CaD-immunostained image shown in A; arrows indicate growth cones. C, The neurons, which had been transfected with GFP as a cell volume maker, were fixed and triple-stained with anti-CaD, anti-GFP, and anti-tau1 antibodies. Bar, 50 µm. D, Dendrite and axon fluorescence intensities were measured in the CaD-immunostained and GFP-immunostained images shown in C, respectively, and then GFP-intensity was subtracted from CaD-intensity to correct for the influence of cell volume. Arrows indicate growth cones. E, Changes in CaD protein expression during neuronal development. F, The morphology of GFP- or GFP-CaD-transfected neurons after 3 days in culture (bar, 100 µm) and
Caldesmon regulates axon extension

G, quantification of their axonal and dendritic length. Axonal length represents the longest axon branch. Data are means ± SE from six independent experiments. H, The morphology of CaD-induced filopodia-like protrusions. The myc-CaD-transfected neurons were fixed and stained with anti-tubulin and phalloidin (F-actin). Bar, 25 μm. I, The morphology of neurons transfected with control miRNA, CaD miRNA1, or CaD miRNA2, and cultured for 5 days (bar, 100 μm), and J, quantification their axonal and dendritic lengths. Data are means ± SE from four to six independent experiments.

FIGURE 2. CaD involvement in cortical NPCs’ axon development. A, Proliferating cortical NPCs were incubated with control siRNA, CaD siRNA1, or CaD siRNA2 for 3 days, and cultured in basic FGF-free medium for 3 days. The cultured cells were fixed and stained with anti-tau1 (green) and anti-MAP2 (red) antibodies. Bar, 200 μm. B, The percentage of tau1-negative non-polarized cells among the differentiating progenitor cells. Data are means ± SE from at least 120 cells. C, Quantification of axonal length in tau-1-positive cells. Data are means ± SE from at least 30 cells.

FIGURE 3. The effect of CaD N- and C-terminal fragments on axon extension. A, CaD, myosin IIA, and myosin IIB localization in the hippocampal neuron axonal growth cone. Bar, 10 μm. B, CaD domain structure. C, Morphology of neurons cultured for 3 days after transfection with GFP, GFP-N-CaD, GFP-C-CaD, or GFP-N-CaD Δ21-47 (bar, 100 μm.), and D, quantification of their axonal length. Data are means ± SE of values from four to six independent experiments. E, quantification of their filipodial density. Data are means ± SE of values from four to six independent experiments.

FIGURE 4. CaD and myosin II interactions. A, HEK 293T cells transfected with myc-CaD and FLAG-HMM IIA or FLAG-HMM IIB were immunoprecipitated using anti-myc antibody or B, anti-FLAG antibody. C, HEK 293T cells were transfected with FLAG-HMM IIA and GFP, GFP-CaD, GFP-N-CaD, GFP-N-CaD Δ21-47, or GFP-C-CaD and immunoprecipitated with anti-GFP antibody. Arrowhead: IgG light chain position. D, HEK 293T cells were transfected with FLAG-HMM IIA and GFP, GFP-CaD Δ21-47, or GFP-CaD and immunoprecipitated with anti-GFP antibody. Arrowhead: IgG light chain position.

FIGURE 5. CaD fragments ectopically expressed in A549 cells. A549 cells transfected with GFP, GFP-C-CaD, GFP-N-CaD, or GFP-N-CaD Δ21-47 were fixed and stained with anti-GFP antibody (green) and phalloidin (F-actin, red). At right, cells were treated with 10 μM blebbistatin (blebb) for 30 min and then fixed and stained with phalloidin. Bar, 100 μm. B, quantification of fluorescence intensity of phalloidin staining in GFP-positive and GFP-negative cells, respectively.
FIGURE 6. The effect of CaD fragments on growth cone morphology and myosin II localization.
Hippocampal neurons transfected with GFP, GFP-CaD, GFP-N-CaD, GFP-N-CaD Δ21-47, or GFP-CaD were fixed and stained with anti-GFP (green) and anti-myosin IIA or IIB (red) antibodies. Bar, 10 µm.

FIGURE 7. Myosin II functions in CaD-enhanced axon extension. A, Hippocampal neurons were cultured with 10 µM blebbistatin, 50 µM ML-7, or 10 µM Y27632 for 3 days, and then fixed and stained with anti-tau1 (green) and anti-MAP2 (red) antibodies (bar, 400 µm.); B, Quantification of axonal length in these neurons. Data are means ± SE of values from at least 70 cells. C, Primary cultured hippocampal neurons transfected with GFP or GFP-CaD and incubated with or without 10 µM blebbistatin for 3 days (bar, 100 µm); D, Quantification of axonal length in these neurons. Data are means ± SE from four independent experiments. E, Growth cone morphology in GFP- or GFP-CaD-transfected neurons before or after treatment with 10 µM blebbistatin for 30 min. To visualize F-actin in living neurons, the cells were transfected with mcherry-LifeAct. Bar, 20 µm.
Fig. 1
Fig. 2
Fig. 3
Fig 4
Fig. 5
Fig. 6
Fig. 7