Filopodial Calcium Transients Promote Substrate-Dependent Growth Cone Turning

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Filopodia that extend from neuronal growth cones sample the environment for extracellular guidance cues, but the signals they transmit to growth cones are unknown. Filopodia were observed generating localized transient elevations of intracellular calcium \([Ca^{2+}]_i\) that propagate back to the growth cone and stimulate global \(Ca^{2+}\) elevations. The frequency of filopodial \(Ca^{2+}\) transients was substrate-dependent and may be due in part to influx of \(Ca^{2+}\) through channels activated by integrin receptors. These transients slowed neurite outgrowth by reducing filopodial motility and promoted turning when stimulated differentially within filopodia on one side of the growth cone. These rapid signals appear to serve both as autonomous regulators of filopodial movement and as frequency-coded signals integrated within the growth cone and could be a common signaling process for many motile cells.

Growth cone filopodia are sensory/motor protrusions that first detect changes in the molecular milieu during axon pathfinding (1–3). Activation of receptors at their tips by interaction with ligands may generate signals that are transmitted back to the growth cone where they are transduced into changes in motility, but both the identity of these signals and whether they regulate filopodial movement are unknown. Although spontaneous changes in \([Ca^{2+}]_i\) within growth cones have been imaged at 0.1 Hz (4, 5), here we analyzed fast \(Ca^{2+}\) dynamics within individual filopodia of cultured Xenopus spinal neurons (6) by high-speed (8 Hz) confocal imaging of magnified regions of Fluo-4 loaded growth cones (7). Most filopodia (55 ± 14%) of neurons grown on poly-D-lysine (PD/L) generated brief \(Ca^{2+}\) transients (317 ± 85 ms) during 30-s imaging periods (Fig. 1, A and B) (n = 22 filopodia from seven growth cones); active filopodia generated transients at 9.2 ± 1 min−1. \(Ca^{2+}\) transients detected by ratiometric imaging of Fluo-4 and Fura-Red fluorescent signals (n = 9) (8, 9) (Fig. 1, C and D) (movie 1, 10) ruled out artifactual fluorescence elevations due to movement or volume changes. \(Ca^{2+}\) elevations occurred locally at the tip or along the length of a filopodium and spread to the growth cone with no apparent decrement at 12.5 ± 0.6 μm/s (n = 12) (Fig. 1, E and F) (movie 2, 10), consistent with either propagated surface entry or intracellular diffusion in a weakly buffered environment (11). Filopodial transients appeared similar to growth cone transients (4, 5) that they depended on \(Ca^{2+}\) influx through nonvoltage-gated channels, but they were distinct because they were not amplified by \(Ca^{2+}\) release from internal stores (Fig. 1G).

To determine the dependence of fast \(Ca^{2+}\) transients on molecules that regulate neurite outgrowth, \([Ca^{2+}]_i\) was imaged in filopodia of neurons grown on eight different substrates (12). Transient frequency varied from one substrate to another whereas duration and amplitude were similar (Fig. 2A) (Web table 1, 10), indicating that these \(Ca^{2+}\) elevations could provide a specific signaling mechanism in growth cone filopodia. Growth cones did not generate global \(Ca^{2+}\) transients on tenascin-C (TN), fibronectin (FN), or vitronectin (VN), indicating that on certain substrates local filopodial transients are not sufficient to produce global elevations (Web table 1, 10). In physiological extracellular \(Ca^{2+}\), the frequency of fast local transients was highest on TN and declined as the TN concentration ([TN]) was lowered (Fig. 2A inset), suggesting that TN stimulates transient production.

Because extracellular matrix (ECM) components, cell adhesion molecules, and artificial substrates stimulate \(Ca^{2+}\) transients in filopodia, \(Ca^{2+}\) entry could be activated by multiple receptor types. To determine whether a particular class of receptor is sufficient to generate these fast \(Ca^{2+}\) transients, we examined the role of β1 integrin receptors that bind many ECM proteins and trigger \(Ca^{2+}\) fluctuations (7, 13). The tripeptide sequence arginine-glycine-aspartate (RGD), contained in many ECM molecules including TN (14) and recognized by a subset of integrins (15), is sufficient to stimulate adhesion and \(Ca^{2+}\) influx in non-neuronal cells (16). Soluble RGDS (S, serine) elevated \([Ca^{2+}]_i\) in growth cones (Fig. 2B) and increased mean filopodial \(Ca^{2+}\) transient frequency from 2.8 ± 0.4 to 4.3 ± 0.8 min−1 (P < 0.05; n = 26 active and inactive filopodia from seven growth cones). The effects of RGDS were specific because the non–integrin binding peptide RGES (E, glutamate) neither elevated \([Ca^{2+}]_i\), nor altered filopodial transient frequency and antibodies that block the function of β1 integrins (17) prevented RGDS-mediated \(Ca^{2+}\) elevations. In addition to its effects on the frequency of filopodial \(Ca^{2+}\) transients, RGDS evoked transients involving larger regions in the growth cone vein (36%, n = 11) (Fig. 2, C and D) (movie 3, 10). These findings show that engagement of integrins on neuronal growth cones can activate \(Ca^{2+}\) influx and oscillations and suggest that integrin clusters (18, 19) may be the sites of fast local \(Ca^{2+}\) transients.

Integrins promote \(Ca^{2+}\) influx and adhesion, which lead to increased cell attachment, spreading, and migration (15). To determine whether the frequency of fast local \(Ca^{2+}\) transients regulates filopodial motility, we measured the lifetime from initiation to retraction of filopodia (20, 21) of growth cones growing on three concentrations of TN that elicit varying frequencies of \(Ca^{2+}\) transients (Fig. 2A inset). As the [TN] increased, the average filopodial lifetime also increased, whereas the average axon length decreased. The effect of TN is \(Ca^{2+}\)-dependent because blocking local transients with the \(Ca^{2+}\) chelator BAPTA reduced filopodial lifetimes (Fig. 3A). Regulation of filopodial lifetime by \(Ca^{2+}\) transients could be due to effects on the actin cytoskeleton (22) or adherence through increased integrin affinity or clustering (avidity) (13). To test whether \(Ca^{2+}\) transients regulate integrin avidity, β1 integrin clustering was examined before and after chelation of \(Ca^{2+}\) with BAPTA during the time when filopodial lifetimes are reduced (23, 24). Blocking endogenous filopodial \(Ca^{2+}\) transients with BAPTA had no detectable effect on integrin clustering at filopodial tips (Fig. 3, B and C). However, localized \(Ca^{2+}\) transients were consistently initiated at sites of punctate integrin clusters (15/15 clusters from 4 growth cones) (Fig. 3, D and E) (Web fig. 1, 10), suggesting that they stabilize filopodia through integrin activation.

To determine whether \(Ca^{2+}\) transients generated in filopodia promote global growth cone transients, we exploited differences in transient frequencies evoked by laminin (LN) and tissue culture plastic (TC). Because growth cones exhibit a higher frequency of

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both local and global Ca\textsuperscript{2+} transients on TC than LN, we investigated the distance at which Ca\textsuperscript{2+} dynamics change as growth cones encounter a LN-TC border (25, 26). The frequency of Ca\textsuperscript{2+} transients in growth cones on LN increased as they approached TC, suggesting that filopodial contact with TC is sufficient to induce global Ca\textsuperscript{2+} changes. The signals promoting global Ca\textsuperscript{2+} transients may be local Ca\textsuperscript{2+} transients because these elevations occurred at a higher frequency within individual filopodia that project onto TC compared with LN (27). To investigate whether filopodial input stimulates the production of global Ca\textsuperscript{2+} transients, global transient frequency was measured in growth cones at this boundary in the presence of 100 nM cytochalasin D to disrupt actin filaments. In the absence of filopodia, the frequency of these transients did not increase until growth cones were within 5 μm of the border (Web fig. 2A, 10). Because Ca\textsuperscript{2+} transients precede growth cone turning in vivo (28), we analyzed whether they induce axon turning at a LN-TC border in vitro. Most axons turned to remain on LN when transients were allowed, where-

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**Fig. 1.** Spontaneous filopodial Ca\textsuperscript{2+} transients visualized by high-speed imaging of Fluo-4 fluorescence. (A) Time-lapse pseudocolor confocal images of Fluo-4 loaded filopodia. Regions selected for quantification at the tip, middle, and base of one filopodium are shown at 0 ms. A local Ca\textsuperscript{2+} transient at the tip occurs at 125 ms (arrow). Fluorescence intensity is coded on a linear pseudocolor scale. (B) Normalized fluorescence changes within regions of this filopodium captured at 125-ms intervals. Ca\textsuperscript{2+} transients occur at high frequency at the tip and move proximally toward the growth cone. Arrowhead indicates transient visualized in (A). F/F\textsubscript{0} indicates Fluo-4 fluorescence relative to baseline fluorescence F\textsubscript{0}. (C) Pseudocolor ratio images of filopodia loaded with Fluo-4 and Fura-Red. (D) Fluorescence ratio increases within regions of the filopodium in (C) demonstrate that these transients are not due to artifacts of movement or volume changes. Note that elevated Ca\textsuperscript{2+}initiated at the tip causes a small elevation of Ca\textsuperscript{2+} at the distal region of this growth cone (base). Arrowhead indicates transient shown in (C). (E) Fluorescence profile images that show elevated Ca\textsuperscript{2+} (colored arrows) moves from tip to base of this filopodium within 375 ms. (F) Fluorescence along the filopodium in (E) normalized to resting levels (lincr scan between white arrowheads in 0-ms panel in (E)) and smoothed with a five-point moving average. The velocity of propagation is 19.2 μm/s. Synchronous fluctuations are artifacts due to scanning at high zoom. (G) Ca\textsuperscript{2+} transients in filopodia on tissue culture plastic are generated by Ca\textsuperscript{2+} influx through a non-voltage-gated channel and are not affected by store release. Ca\textsuperscript{2+} influx is required because the average transient frequency decreased in reduced [Ca\textsuperscript{2+}]\textsubscript{o}, and transients were abolished in 0 mM Ca\textsuperscript{2+}, with 10 mM NiCl in the presence of 2 mM Ca\textsuperscript{2+} or by intracellular Ca\textsuperscript{2+} chelation with BAPTA. Filopodial transient frequency was unaffected by voltage-gated Ca\textsuperscript{2+} channel blockers (VGC) (50 μM NiCl, 10⁻⁷ M nifedipine, 10⁻⁶ M Ω-conotoxin, and 60 nM Ω-agatoxin) or Ca\textsuperscript{2+} store release with thapsigargin (TG). Neurons were grown on poly-d-lysine (PDL) in 2 mM Ca\textsuperscript{2+} except where indicated and imaged at 12 to 24 hours in culture. n > 20 filopodia for each condition. Bar in (A), (C) and (E), 2 μm.
Ca²⁺ and veil transients. Antibodies to traces are averages from 10 growth cones and stimulated increased frequency of ßlopodial Ca²⁺.

Normalized ßuorescence changes within the active veil in (C) captured at 125-ms intervals.

Inactive ones) of neurons grown in 2 mM Ca²⁺ through integrin receptors. (Web fig. 2B, I0). These results indicate that when ßlopodia contact a second substrate, local Ca²⁺ signals spread to the growth cone and stimulate global Ca²⁺ elevations that promote growth cone turning.

To test whether ßlopodial Ca²⁺ transients on one side of the growth cone are sufficient to promote turning, we examined growth cones at a boundary between high (50 μg/ml) and low (5 μg/ml) TN (25). Because growth cones do not generate global Ca²⁺ elevations on TN, Ca²⁺-dependent turning at a high [TN]/low [TN] border would likely result from differences in the frequency of ßlopodial Ca²⁺ transients on varying [TN] (Fig. 2A).

Fig. 3. Local Ca²⁺ transients reduce ßlopodial motility and slow neurite outgrowth. (A) Increased [TN] prolonged ßlopodial lifetimes and shortened neurite lengths. The effect of [TN] depended on Ca²⁺ transients because 100-nM BAPTA ([TN]) did not block spontaneous ßlopodial Ca²⁺ transients, perhaps due to restricted antibody access. (C) Pseudocolor proﬁle images show that elevated Ca²⁺ (arrow) in response to 1 mM soluble RGDS spreads through a growth cone veil in 1 s. Bar, 2 μm. Fluorescence intensity is coded on a linear pseudocolor scale. (D) Normalized fluorescence changes within the active veil in (C) captured at 125-ms intervals. Ca²⁺ transients occurred at high frequency after the addition of RGDS (arrow). Gap in trace represents 14 s during application. Neurons in (B) through (D) were grown on TN (5 μg/ml) in 2 mM Ca²⁺.

Fig. 2. Frequencies of local ßlopodial Ca²⁺ transients are regulated by the substrate, in part through integrin receptors. (A) The average frequency of transients for all ßlopodia (including inactive ones) of neurons grown in 2 mM Ca²⁺ on artiﬁcial substrates (TC, PDL), cell adhesion molecules [N-cadherin (Cad) and L1], and on ECM components LN, FN, TN, and VN (50 μg/ml). For some substrates (L1, Cad, LN, PDL, and TC) the frequency of global growth cone transients correlated with the local ßlopodial transient frequency, but no correlation was seen for others (FN, VN, TN). TN stimulated the highest frequency of transients in ßlopodia, in a dose-dependent manner (inset). n = 10 ßlopodia for each condition. (B) Soluble RGDS produced a dose-dependent, biphasic elevation of [Ca²⁺], in 75% of growth cones tested (arrowhead; traces are averages from 10 growth cones) and stimulated increased frequency of ßlopodial and veil transients. Antibodies to β1 integrin did not block spontaneous ßlopodial Ca²⁺ transients, perhaps due to restricted antibody access. (C) Pseudocolor proﬁle images show that elevated Ca²⁺ (arrow) in response to 1 mM soluble RGDS spreads through a growth cone veil in 1 s. Bar, 2 μm. Fluorescence intensity is coded on a linear pseudocolor scale. (D) Normalized ßuorescence changes within the active veil in (C) captured at 125-ms intervals. Ca²⁺ transients occurred at high frequency after the addition of RGDS (arrow). Gap in trace represents 14 s during application. Neurons in (B) through (D) were grown on TN (5 μg/ml) in 2 mM Ca²⁺.

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Thus a step-gradient of TN directs axon growth in a Ca²⁺-dependent manner by stimulating ßlopodial Ca²⁺ transients on one side of the growth cone.

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Because different concentrations of TN may affect axon growth in ways unrelated to filopodial Ca\(^{2+}\) transients, we locally released caged-Ca\(^{2+}\) within filopodia on a homogeneous substrate to test whether local differences in filopodial transient frequencies are sufficient to produce turning (29). Growth cones loaded with caged-Ca\(^{2+}\) (NP-EGTA) or unloaded controls were positioned so their forward projecting filopodia extended into a 20-μm spot that was pulsed with UV light (310 to 410 nm) for 200 ms at 10-s intervals. Because nonphotolyzed NP-EGTA chelates spontaneous transients due to its low dissociation constant (\(K_D\)) (80 nM) (30), Ca\(^{2+}\) transients were generated only in those filopodia projecting into the spot (Fig. 4, C and D). After 5 to 10 μm of growth, the direction of axon outgrowth was deflected by 37° ± 8° (n = 10) in all growth cones loaded with caged-Ca\(^{2+}\), but only by 14° ± 3° (n = 10) in unloaded controls (Fig. 4, E and K) \((P < 0.02)\). These results show that filopodial Ca\(^{2+}\) transients imposed on one side of the growth cone are sufficient to induce turning away from that side.

Our findings indicate that growth cones exhibit spontaneous cytosolic Ca\(^{2+}\) signals over a wide range of spatial and temporal domains that have diverse effects on pathfinding behaviors. The amplitudes of Ca\(^{2+}\) gradients imposed within the palm of growth cones also regulate attractive or repulsive turning (31, 32). The frequency of Ca\(^{2+}\) transients in filopodia depends on substrate type and concentration, underscoring their biological significance. Ca\(^{2+}\) elevations detected in isolated filopodia in contact with guidance cues may be initial signaling events necessary for adhesion (33). Integrin-mediated Ca\(^{2+}\) influx through non–voltage-gated channels seems to contribute to Ca\(^{2+}\) signals that result in decreased filopodial motility, possibly by feedback activation of integrins (33). Although integrins appear to be sufficient to stimulate Ca\(^{2+}\) transients, they are not necessarily given activation by non–integrin binding substrates. Filopodial Ca\(^{2+}\) transients are similar to other types of elemental signaling processes such as vesicular transmitter release (34) and Ca\(^{2+}\) puffs and sparks (35), because they can be summed to produce global actions on certain substrates. The autonomous effects of local Ca\(^{2+}\) signals on filopodial motility and substrate-specific ability to induce global elevations of [Ca\(^{2+}\)]\(_i\), together provide versatile control of growth cone behavior.

References and Notes
7. For Ca\(^{2+}\) imaging, neurons were loaded with 2 to 5 μM Fluo 4-AM (with 0.01% pluronic acid/0.1% dimethyl sulfoxide (DMSO) in modified Ringer’s solution (MR) [3]. Molecular Probes] for 1 hour at 20°C, rinsed, and imaged at frequencies of 0.06 to 8 Hz on Bio-Rad MRC 600 argon laser or MRC 1024 krypton/argon laser confocal systems. Eight-Hz imaging was performed with the MRC 1024 mounted on an Olympus BX70 upright microscope.
microscope using either a 60× water (NA 0.9) or 100× oil (NA 1.4) objective. An acquisition box of 100 pixels by 40 pixels was used to increase collection speed and zoomed to achieve pixel sizes between 0.2 and 0.4 μm. Reduced speed Ca2⁺ imaging (0.06 to 1 Hz) used for RGD experiments (RGDS and RCGE from Sigma) and imaging of global growth cone transients was performed on the MRC 600 mounted on a Zeiss upright microscope using a 764 pixel by 512 pixel collection box and a 40× Fluor (NA 1.2) objective. For all experiments, laser power was attenuated to 1 to 10% of maximum using neutral density filters. Loose coverslips with adherent neurons were inverted for use with oil objectives by first sealing them on a coverside with high-vacuum grease (Dow Corning). Inlet and outlet ports were left when necessary for exchange of solution.

9. For combined Fluo-4/Fura-Red imaging, neurons loaded with 5 μM Fluo-4-AM and 5 μM Fura-Red-AM (in 0.01% pluronic acid) were excited with the 488-nm laser line of the MRC 1024 and images were collected at both 522 ± 35 and at ±855 nm wavelength. Fluorescence was quantified and changes were normalized using MetaMorph (Universal Imaging) or NIH Image (W. Rasband, NIH) software; events exceeding 10% of baseline fluorescence were scored as filopodial transients. Statistical significance was determined using Student’s t-test unless otherwise noted, and variance is reported ± SEM.

10. Web figures 1 and 2, Web table 1, and three Quicktime movies are available at Science Online at www.sciencemag.org/cgi/content/full/291/5510/1983/DC1.
12. Cultures of dissociated spinal neurons and neural tube explants were prepared from stage 20 to 22 Xenopus embryos. Cells were grown at 20°C in MR containing 2 mM Ca²⁺ (3) on untreated tissue culture plastic or on various biological substrates coated onto tissue culture plastic or acid-washed glass coverslips, usually at saturating concentrations [LN, FN, VN, and Poly-D-lysine from Sigma; TN from Gibco].
21. For analysis of filopodial lifetimes and for Ca⁺⁺ photorelease studies, neurons were labeled with green fluorescence protein-actin (GFP-actin) to enhance visualization of filopodia. Two individual blastomeres at the eight-cell stage were injected with 1 to 5 ng of capped GFP-actin mRNA (mMessage machine, Ambion). Filopodial lifetime measurements were made on growth cones from loaded (1 hour, 100 μM BAPTA AM; Molecular Probes), and unloaded cultures on 5, 10, or 50 μg/ml TN. Motile growth cones were scored and imaged at 15-s intervals for 30 min with the MRC 600 using a 100× objective. A programmable shutter (Uniblitz) pulsed this spot with 310- to 410-nm ultra-violet light filtered for a 100-W mercury lamp for a duration of 200 ms every 10 s. The direction of neurite growth was measured from the center of the leading edge to middle of the neurite at the base of the growth cone using NIH Image software. The maximum difference in the angle of neurite growth between 5 to 15 μm of growth was scored. Neurites with a net extension of <5 μm over the 30-min period were not included. Photorelease of Ca²⁺ in selected filopodia was tested using non–GFP expressing neurons loaded with 2.5 μM Fluo-4 and NP-EGTA.
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Requirement for the SLP-76 Adaptor GADS in T Cell Development

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GADS is an adaptor protein implicated in CD3 signaling because of its ability to link SLP-76 to LAT. A GADS-deficient mouse was generated by gene targeting, and the function of GADS in T cell development and activation was examined. GADS-CD4⁺CD8⁺ thymocytes exhibited a severe block in proliferation but still differentiated into mature T cells. GADS⁺ thymocytes failed to respond to CD3 cross-linking in vivo and were impaired in positive and negative selection. Immunoprecipitation experiments revealed that the association between SLP-76 and LAT was uncoupled in GADS⁺ thymocytes. These observations indicate that GADS is a critical adaptor for CD3 signaling.

The development and function of T cells are regulated by signaling through the CD3 complex, which serves both the pre-T cell receptor (pre-TCR) and the TCR (β/γ) and references therein). Cross-linking of CD3 induces protein tyrosine phosphorylation in a wide range of proteins. Among these phosphorylation targets are two adaptor proteins, LAT and SLP-76, which function in a coordinated fashion to activate a diverse set of signaling proteins (2–5). The critical function of SLP-76 and LAT is supported by the observation that mice lacking SLP-76 or LAT exhibit an absolute block in early thymocyte development (6–8).

The function of SLP-76 is dependent on its association with LAT (9–13). This association is mediated by an adaptor known as GADS, which contains two SH3 domains flanking a SH2 domain and a linker region. GADS associates constitutively with SLP-76 through the binding of the GADS SH3 domain, and is recruited to LAT through binding of its SH2 domain to phosphorysine motifs on LAT upon TCR activation (9–13). Besides GADS, Grb2 and possibly Grb7 are also implicated as adaptors for SLP-76 (4). Because mutant T cells or primary mast cells lacking LAT demonstrate reduced phosphorylation of SLP-76 upon receptor activation.