Protein Synthesis in Distal Axons Is Not Required for Growth Cone Responses to Guidance Cues

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Recent evidence suggests that growth cone responses to guidance cues require local protein synthesis. Using chick neurons, we investigated whether protein synthesis is required for growth cones of several types to respond to guidance cues. First, we found that global inhibition of protein synthesis stops axonal elongation after 2 h. When protein synthesis inhibitors were added 15 min before adding guidance cues, we found no changes in the typical responses of retinal, sensory, and sympathetic growth cones. In the presence of cycloheximide or anisomycin, ephrin-A2, slit-3, and semaphorin3A still induced growth cone collapse and loss of actin filaments, nerve growth factor (NGF) and neurotrophin-3 still induced growth cone protrusion and increased filamentous actin, and sensory growth cones turned toward an NGF source. In compartmented chambers that separated perikarya from axons, axons grew for 24 – 48 h in the presence of cycloheximide and responded to negative and positive cues. Our results indicate that protein synthesis is not strictly required in the mechanisms for growth cone responses to many guidance cues. Differences between our results and other studies may exist because of different cellular metabolic levels in in vitro conditions and a difference in when axonal functions become dependent on local protein synthesis.

Key words: growth cone; guidance; protein synthesis; axon; actin; RhoA

Introduction

Neurons are highly polarized with elaborate shapes and highly localized activities. Protein synthesis is not confined to the perikaryon; rather, proteins made in dendrites and axons have important functions (Steward and Schuman, 2003). Defective dendritic protein translation underlies some behavioral deficits and mental retardation (Bramham and Wells, 2007; Dahm and Macchi, 2007; Wang et al., 2007, Axonal protein synthesis is critical in functions of terminals, as well as in development and regeneration (Koenig and Giuditta, 1999; Campenot and Eng, 2000; Zhang and Poo, 2002; Piper and Holt, 2004; Verma et al., 2005; Twiss and van Minnen, 2006; Hengst and Jaffrey, 2007).

Is local protein synthesis important in axonal growth and guidance? In vitro experiments with compartmented chambers indicate that axonal protein synthesis is not required for axonal growth (Eng et al., 1999; Blackmore and Letourneau, 2007), yet there are reports that responses to guidance cues require axonal protein synthesis (Campbell and Holt, 2001; Guirland et al., 2003; Brunet et al., 2005; Wu et al., 2005; Leung et al., 2006; Piper et al., 2006; Yao et al., 2006; Lin and Holt, 2007; Farrar and Spencer, 2008). When growth cones enter new environments, local synthesis of receptors could sensitize growth cones to new guidance cues (Brittis et al., 2002). In addition, growth cone adaptations to changed balances of cues may involve local synthesis of receptors or signaling components (Ming et al., 2002).

Thus, local synthesis of receptors or signaling components may be necessary for growth cone navigation. Is it also necessary to locally make proteins with general roles in growth cone motility? Recent papers report that RhoA GTPase and β-actin must be rapidly and locally synthesized for growth cones to respond to several cues (Wu et al., 2005; Leung et al., 2006; Yao et al., 2006). In light of the diverse mechanisms that rapidly regulate RhoA and β-actin activity, it is unexpected that these components must be locally synthesized to enable responses to guidance cues.

Because previous studies involved few neuronal types, we investigated whether protein synthesis is required for growth cones of other neurons to respond to guidance cues. First, we found that globally inhibiting protein synthesis leads to slowing and arrest of axonal growth over several hours. When protein synthesis inhibitors were added before several positive and negative cues, we found normal growth cone responses. Repulsive cues, ephrin-A2, slit-3, Semaphorin3A (Sema3A), and Semaphorin6A (Sema6A), induced growth cone collapse in the presence of protein synthesis inhibitors. Filamentous actin (F-actin) polymer was lost after adding repulsive cues. Furthermore, attractive cues, nerve growth factor (NGF) and neurotrophin-3 (NT-3), induced growth cone protrusion and increased F-actin despite inhibiting protein synthesis. In addition, growth cones turned toward an NGF source in the presence of protein synthesis inhibitors. In compartmented cultures in which axonal protein synthesis was inhibited for many h, growth cones still responded to guidance cues. Our data indicate that protein synthesis in distal axons of chick retinal, sympathetic, and dorsal root ganglia (DRG) neurons and mouse DRG neurons is not required for responses to several guidance cues.
Materials and Methods

**Materials.** F-12 medium, DMEM medium, B27 additives, laminin, poly-D-lysine, Alexa Fluor 488–phalloidin, Alexa Fluor 562 secondary antibodies, Click-iT AHA (catalog #C10102) and Click-iT Tetramethylrhodamine Protein Analysis Detection kit (catalog #C33370) reagents were purchased from Invitrogen. NGF, NT-3, L1–Fc, slit-3, ephrin-A2–Fc, Sema3A–Fc, and Sema6A–Fc were purchased from R & D Systems. Anti-RhoA monoclonal antibodies were purchased from Santa Cruz Biotechnology, and anti-eliaF-4EBP-P rabbit antibodies were purchased from Cell Signaling Technology. Rabbit anti-b-actin was a gift from Dr. James Ervasti (University of Minnesota, Minneapolis, MN) (Prins et al., 2008). Collagen was purchased from Inamed. Cycloheximide and anisomycin were purchased from Calbiochem, and M&G-132 (N-(phenylmethoxy)carbonyl)-s-leucyl-N-((1S)-1-formyl-3-methylbutyl)-s-leucinamide) was from Tocris Bioscience.

**Neuronal culture.** Culture dishes, glass coverslips, or 24- or 96-well dishes were coated overnight with 20 µg/ml laminin or 4 µg/ml recombinant L1–Fc mixed with 8 µg/ml Fc in PBS. Explants of embryonic day 7 (E7) and E13 DRGs, E7 temporal retina, and E7 sympathetic ganglia were dissected from chick embryos, and DRGs were removed from E15 mouse embryos, according to procedures approved by the University of Minnesota Institutional Animal Care and Use Committee. Neural tissues were cultured on experimental substrates in F-12 with B27 additives and buffered to pH 7.4 with 10 mM HEPES. Neurotrophins, NGF or NT-3, were added to cultures, as noted. Neural tissues were cultured overnight in a humidified incubator at 37°C.

**Neuronal transfection.** E7 chick DRGs were dissected and dissociated with 0.2% bacto-trypsin in calcium- and magnesium-free (CMF) PBS for 10 min and then rinsed with F-12H containing 10% serum. Approximately 2 x 10^5 cells were transfected with 3 µg of a plasmid encoding the fluorescent F-actin probe, green fluorescent protein (GFP)–urokinase (UckH) kindly provided by Dr. W. M. Bement (University of Wisconsin, Madison, WI) (Burkel et al., 2007), using the G-13 program of the Amaha Biosystems Nucleofector. Cells were then rinsed with F-12H and cultured overnight in F-12 with B27 additives on video dishes coated with poly-D-lysine, nitrocellulose, and L1.

**Axon elongation assay.** The effect of protein synthesis inhibitors on axon elongation was determined by time-lapse microscopy of axon elongation by DRG or retinal explants cultured in F-12 with B27 (and 10 ng/ml NGF for DRGs) on laminin- or L1-coated 96-well dishes. A dish was placed on the warmed stage of a Nikon inverted microscope, and a microscope field of axons was chosen. Cycloheximide (CHI) or anisomycin (or PBS for control) was added at 20 µM, and time-lapse images were collected at 2–5 min intervals for 2–5 h. Axon elongation rates of sample populations were determined, using MetaMorph software (Molecular Devices), to measure the distance a growth cone migrated in 5–10 min intervals over a 3–5 h period.

**Growth cone collapse assay.** Collapse assays were conducted in 24- or 96-well dishes. Explants were cultured overnight. NGF, 10 ng/ml, was added to cultures of DRGs or sympathetic ganglia. CHI or anisomycin was added in PBS at 20 µM for 15 min. Control dishes received PBS. After 15 min guidance molecules, Sema3A (0.5 µg/ml), ephrin-A2 (1 µM/ml), or slit-3 (3 µg/ml) were added for 30 min in the continued presence of protein synthesis inhibitors. Control dishes received PBS. After 30 min, explants were fixed with 0.5% glutaraldehyde in PBS for 20 min at 37°C. After rinsing in PBS, the wells were visualized with a 20x phase-contrast objective, and growth cones were counted as collapsed if they had one or no filopodia.

**Growth cone spreading assay.** DRG explants were cultured overnight in 96-well dishes. A dish was placed on a warmed stage of an inverted microscope, a field of axons was chosen, and images were collected once per minute for 15 min. Then, 40 ng/ml NGF or NT-3 was added, and images of the same field were collected every minute for 30 min. Growth cone spreading was measured using MetaMorph software to outline the borders of growth cones and determine the enclosed areas.

**Growth cone turning assay.** DRG explants were cultured overnight in F-12 with B27 on laminin-coated coverslips glued over a hole in the bottom of a culture dish. Micropipette tips were dipped in a 1% nitrocellulose solution and dried. After drying, a tip was dipped several times in a solution of 1 µg/ml NGF or BSA in PBS. The micropipette was mounted on a micromanipulator. A culture dish was placed on the warmed microscope stage, and the micropipette tip was positioned 50 µm from a growth cone at a 45° angle to the direction of axon elongation. Images were acquired at 30 s intervals for 15 min before and 45 min after introducing the micropipette tip. CHI was added to some dishes at the beginning of image acquisition. Growth cone turning angles were determined as the change in direction of growth cone migration between the beginning and end of the image acquisition period (Ming et al., 1997).

**Compartmented cultures.** Compartmented cultures were made by putting an E7 DRG explant with attached peripheral root on a substrate and adhering a 5 mm glass cloning cylinder to the substrate with silicone grease, so the DRG was within the lumen and the peripheral root extended under the 1 mm cylinder wall with its terminus outside the wall. This created two compartments and allowed us to expose the DRG and the peripheral root to different media. For chemotaxis experiments, a 3 mg/ml collagen gel was created in the outside compartment, following the instructions of the manufacturer (PurCol; Invitrogen). Affi-Gel Blue Gel chromatography beads were soaked in 1 µg/ml NGF or BSA in PBS for 1 h, rinsed in PBS, and then placed in the collagen gel near the peripheral nerve root with an NGF bead on one side and a BSA bead on the other side of the root.

**Immunocytochemistry.** For immunocytochemistry, explants cultured on laminin- or L1-coated coverslips were fixed with warmed 4% paraformaldehyde in PBS with 5% sucrose at pH 7.4 for 30 min. After rinsing, explants were treated 15 min with 0.1% sodium glycine in PBS and extracted with 0.1% Triton X-100 (TX-100) in PBS with 2% goat serum and 1% BSA for 1 h. Coverslips were incubated with the primary antibodies diluted in PBS containing 1% BSA for 1 h. Primary antibody dilutions were 1:100 for anti-RhoA and anti-eliaF-4EBP-P, and anti-b-actin was used, per instructions from Dr. Ervasti (Prins et al., 2008). For labeling F-actin, Alexa Fluor 488–phalloidin was applied at 2.5 µl/100 µl mixed with the primary antibodies. The coverslips were rinsed and then incubated in 0.1% TX-100 in PBS with 2% goat serum and 1% BSA for 1 h. Then, Alexa Fluor 568 goat anti-rabbit or anti-mouse antibodies at 1:1000 dilution were applied in PBS with 1% BSA for 1 h. After rinsing, the coverslips were incubated with secondary antibody 0.1% TX-100 PBS with 2% goat serum and 1% BSA for 30 min, rinsed, and mounted in anti-fading medium.

**Nascent protein synthesis.** To measure effects of protein synthesis inhibitors CHI and anisomycin on cultured DRG neurons, we used Click-iT reagents from Invitrogen. E7 DRGs were dissociated with 0.2% bacto-trypsin in CMF-PBS for 15 min, rinsed, and plated on poly-D-lysine- and laminin-coated glass coverslips in DMEM with B27 additives and 10 ng/ml NGF. After overnight culture, the dishes were rinsed twice with warmed PBS and incubated in methionine-free DMEM for 45 min. During the last 15 min of this period, 20 µM CHI or anisomycin or control buffer were added to experimental dishes. Then, the medium was replaced with methionine-free DMEM to which 50 µM of the methionine analog l-azidohomoalanine (AHA) (catalog #C10102; Invitrogen) was added. AHA was omitted from some dishes as a negative control. The dishes were returned to the incubator for 60 min for incorporation of the AHA into nascent proteins with or without protein synthesis inhibitors. After 60 min, the dishes were rinsed twice and incubated 30 min in methionine-free DMEM with or without protein synthesis inhibitors. The cultures were fixed with 2% paraformaldehyde and 5% sucrose in PBS for 30 min and then rinsed and incubated in PBS with 0.1% sodium glycine for 15 min. The coverslips were then drained and placed on Parafilm in a humidified 150 mm Petri dish. Using the nomenclature of the Click-iT reagent kit (catalog #C3337; Invitrogen), the carboxytetramethylrhodamine alkyne (TAMRA) reagent was prepared by rapidly mixing 150 µl of the Step 1 solution with 120 µl of distilled water, then 15 µl of Component C was added with 5 s vortexing, followed by 15 µl of Step III solution with 5 s vortexing. Fifty microliters of this solution was applied to each coverslip, followed by 5 µl of the Step 2 solution, which was rapidly mixed with the solution on each coverslip. The coverslips were then incubated 30 min in the dark. Then, the coverslips were drained, rinsed twice with distilled water, then twice with methanol for 15 min,
before being returned to distilled water, and finally mounted on glass slides in anti-fading medium.

To measure the effects of NGF and Sema3A on protein synthesis in DRG neurons, we used the Click-IT reagents as described above. Dissociated DRG neurons were cultured overnight on poly-lysine- and laminin-coated coverslips in DMEM and B27 additives with 0.1 ng/ml NGF. After rinsing and 45 min incubation in methionine-free DMEM, the medium was replaced with methionine-free DMEM containing 50 μM AHA and 40 ng/ml NGF, 1 μg/ml Sema3A, or PBS as a control for addition of a guidance cue. Other samples were incubated in methionine-free DMEM without AHA, as a control for the TAMRA reagent procedure. After 1 h incubation with AHA and the guidance cues, the dishes were rinsed twice and incubated in methionine-free DMEM for 30 min, followed by fixation and incubation with the TAMRA reagents, as described above.

Quantitative fluorescence determinations. A Spot digital camera mounted on an Olympus XC-70 inverted microscope was used to acquire all images for quantitative fluorescence measurements. In any one experiment, all coverslips with explants or cells were fixed and processed together. For collection of fluorescence images, all images for one experiment were acquired in a single session. Parameters of time interval and gain setting on the digital SPOT camera were adjusted so the brightest areas did not reach saturation, and the same gain and time interval was used to capture all images of any particular staining. All analysis of image intensities was conducted from a similar region at the center of the microscope field.

MetaMorph software (Molecular Devices) was used for all image analysis. Software tools were used to determine fluorescence intensity along axons, using a line-scan tool, within growth cones, using a line tool to outline the terminal 25 μm of each distal axon and growth cone, and a square-shaped tool placed within the center of neuronal somata. After acquisition of each fluorescent neuronal image, a background was acquired by moving the software tool slightly outside the boundary of the neuron. The background fluorescence intensity value was subtracted from the fluorescence intensity value of the accompanying neuronal measurement.

Effects of NGF and Sema3A on phospho-eIF-4E-BP content of distal axons. E7 DRG explants were cultured overnight on laminin-coated coverslips. NGF at 40 ng/ml, 1 μg/ml Sema3A, or PBS was added for 15 min, and then the cultures were fixed and processed for immunofluorescence staining, as described above. For measuring anti-eIF-4E-BP-P staining, images of immunofluorescence staining of a sample population of distal axons and growth cones were acquired using a 60× oil-immersion objective. To measure F-actin content, the boundary of each growth cone was outlined with the outline tool, and the mean pixel intensity and integrated pixel intensities of F-actin staining were determined for each growth cone.

Effects of NGF on F-actin content of DRG growth cones. E7 DRG explants were cultured overnight on laminin-coated coverslips. Twenty micromolar CHI or PBS was added for 15 min, and then 40 ng/ml NGF (DRGs) or 2 μg/ml ephrin-A2 (temporal retina) was added for 15 min in the continued presence of CHI or PBS. The cultures were fixed with 4% paraformaldehyde and 5% sucrose for 30 min and then extracted with 0.1% TX-100 in PBS for 15 min. F-actin was stained by incubation with 2.5 μl/0.1 ml Alexa Fluor 568–phalloidin in PBS for 60 min. After rinsing, the coverslips were mounted in anti-fading medium. Images of sample populations of growth cones were acquired with a 60× oil-immersion objective. Using the line-scan tool of MetaMorph, phalloidin fluorescence was determined along a 2-pixel-wide by 6 μm line backward from the center of the growth cone leading margin.

Effects of NGF on F-actin content of DRG growth cones. E7 DRG explants were cultured overnight on laminin-coated coverslips. Twenty micromolar CHI or PBS was added to cultures for 15 min, and then 40 ng/ml NGF was added for 15 min in the continued presence of CHI or PBS. The cultures were fixed with 4% paraformaldehyde and 5% sucrose for 30 min and then extracted with 0.1% TX-100 in PBS for 15 min. F-actin was stained by incubation with 2.5 μl/0.1 ml Alexa Fluor 568–phalloidin in PBS for 60 min. After rinsing, the coverslips were mounted in anti-fading medium. Images of growth cones were acquired with a 60× oil-immersion objective. To measure F-actin content, the boundary of each growth cone was outlined with the outline tool, and the mean pixel intensity and integrated pixel intensities of F-actin staining were determined for each growth cone.

Effects of NGF gradient on GFP–UtrCH distribution in DRG growth cones. Immediately after placing a video dish with GFP–UtrCH-transfected neurons on a warmed microscope stage, 20 μm cycloheximide or the same volume of control media was added for 15 min. A transfected growth cone was identified, and a fluorescent image was taken. An NGF- or BSA-coated micropipette was brought to one side of the growth cone, and after, 2 min, another image was acquired using the same exposure and gain settings. Using MetaMorph software, a 15-pixel-thick line was drawn across the growth cone (perpendicular to its neurite axis) at the two time points before and after applying the NGF gradient. The “line-scan” function was used to obtain fluorescence intensity measurements for each point along the line across the growth cone. Ten growth cones for the control and cycloheximide conditions were used in the analysis.

Statistical analysis. Parameters of population values are reported as mean ± SEM. All statistical analysis was by unpaired Student’s t test or Mann–Whitney U test.

Results

NGF and Sema3A activate protein synthesis in DRG neurons

It is reported that guidance cues trigger inactivation of the translation repressor elf-4EBP1 in distal axons (Campbell and Holt, 2001; Li et al., 2004; Cox et al., 2008). Inactivation of elf-4EBP1 by phosphorylation releases elongation factor elf-4E to initiate mRNA translation. We used phospho-specific antibodies to ask whether the guidance cues NGF and Sema3A induce elf-4EBP1 phosphorylation in axons extended from E7 chick DRG. DRG explants were cultured overnight and exposed to NGF or Sema3A for 15 min before fixation and staining with anti-elf-4EBP1-P (Fig. 1a–c). Figure 1d shows analysis of mean anti-elf-4EBP-P staining of a line scanned from the center of each growth cone proximally for 15 μm in the axon. The mean staining intensity for elf-4EBP1-P was significantly greater in distal axons of both NGF- and Sema3A-treated DRG cultures than in axons of untreated DRGs. Thus, like previous reports, signaling triggered by NGF and Sema3A inhibits a repressor of mRNA translation in distal axons of DRG neurons.

The inactivation of elf-4EBP1 and activation of elongation initiation factor 4E are associated with increased protein synthesis in response to growth factors and guidance cues (Campbell and Holt, 2001; Takei et al., 2001; Li et al., 2004). We investigated whether protein synthesis in DRG neurons is altered by NGF and
Sema3A. To detect nascent proteins, we used the nonradioactive reagent AHA, a methionine analog that is incorporated into proteins instead of methionine and is detected by reaction of the azido-modified protein with a fluorescent alkyne (for details, see Materials and Methods). Overnight cultures of dissociated E7 DRGs were exposed to NGF or Sema3A in AHA-containing media for 60 min before fixation and reaction with the fluorescent alkyne. To control for the labeling procedures, some samples were not given AHA, and, to control for the addition of guidance cues, other samples were given AHA but neither NGF nor Sema3A. As shown in Figure 2a–c, the mean staining intensity for nascent proteins in both cell bodies and distal axons was increased 60% by NGF addition compared with cultures not treated with NGF. After addition of Sema3A, the nascent protein amounts were also elevated from the levels in untreated control neurons. Because this experiment involved one time point, we make no conclusion about effects of NGF or Sema3A on rates of protein synthesis or degradation. Because of the global presentation of ligands and the duration of treatment, we infer nothing about the location of nascent protein synthesis. However, our results indicate that NGF and Sema3A stimulate increased amounts of nascent proteins in cell bodies and axons of DRG neurons. Thus, like previous reports, we found that guidance cues increase nascent proteins in neurons (Zhang et al., 1999, 2001; Campbell and Holt, 2001).

**DRG axonal elongation is inhibited by global treatment with protein synthesis inhibitors but not by inhibiting protein synthesis in distal axons only**

Before assessing the effects of inhibiting protein synthesis on responses to guidance cues, we showed that CHI, which inhibits peptide chain elongation, and anisomycin, which inhibits peptide bond formation, are effective inhibitors of chick neuronal protein synthesis. These drugs were used previously to inhibit protein synthesis in chick neurons (Ludueña, 1973; Oppenheim et al., 1990; Blackmore and Letourneau, 2007). To measure the effects of CHI and anisomycin, we used the same Click-it reagents that we used to determine the effects of NGF and Sema3A on nascent protein synthesis. After washing out methionine-containing DMEM medium, DRG cells were incubated for 1 h with 50 mM AHA with or without 20 μM CHI or anisomycin. As shown in Figure 2f–i and quantitated in j, CHI- and anisomycin-treated neurons were as weakly labeled as control neurons that were not incubated with the methionine analog AHA but only with the fluorescent alkyne reagents. Thus, we could not detect protein synthesis in DRG neurons treated with CHI or anisomycin.

Because it was reported previously that global application of protein synthesis inhibitors did not inhibit axonal extension during a 1 h period (Campbell and Holt, 2001), we investigated axonal elongation by DRG and retinal neurons in the global presence of protein synthesis inhibitors. Explants of E7 DRGs and temporal retina were grown in dishes coated with laminin or the cell adhesion molecule L1. CHI, anisomycin, or the drug vehicle was added, and rates of axonal elongation were calculated after time-lapse imaging. The mean elongation rate for untreated DRG or retinal axons did not change over 4 h of recording, although the elongation rate for DRG axons was significantly faster on laminin than on L1 (130 ± 6 vs 41 ± 6 μm/h, respectively). During the first hour of treatment with 20 μM CHI or anisomycin, the axonal elongation rate of treated DRG or retinal neurons was nearly equal to that of control, untreated neurons (Table 1, Fig. 3a). During the second hour, axonal growth rates began to slow in the presence of protein synthesis inhibitors (Table 1, Fig. 3a). Some growth cones became smaller and less motile, and, on laminin, some axons retracted. On L1, axonal retraction during the second hour of treatment was less frequent than on laminin, but the rate of axon elongation still slowed (Table 1). Continuing after 2 h of protein synthesis inhibition, nearly all axons retracted or became quiescent (Fig. 3b) (supplemental Fig. S1, available at www.jneurosci.org as supplemental material).

We asked whether concurrent inhibition of proteolysis might extend the time that axons elongate in the presence of protein synthesis inhibitors. We added MG-132, an inhibitor of proteosome activity used previously with neurons (Lee and Goldberg 1996; Tursun et al., 2005), to some wells that received CHI. MG-132 alone did not reduce axon elongation over 4 h (Fig. 3b). When CHI was combined with MG-132, fewer axons retracted during hours 2–5 in CHI plus MG-132 than in wells treated with CHI alone, but axonal growth had still stopped in CHI plus MG-132 by 4–5 h (Fig. 3b). Thus, global inhibition of protein synthesis does not significantly inhibit axonal elongation during the first hour of treatment, but, during the second hour, axonal growth slowed and axons began to retract or lose growth cone activity. Concurrent inhibition of protein degradation prolonged the period during which axons could elongate in the absence of protein synthesis.

Previously, compartmented chambers were used to show that protein synthesis in distal axons is not required for axonal elongation (Eng et al., 1999; Blackmore and Letourneau, 2007). We devised a compartmented dish to investigate whether distal protein synthesis is required for DRG axon elongation. An E7, DRG with the attached peripheral root was placed in the lumen of a 5-mm-diameter glass cloning cylinder, which was stuck to the substrate with silicon grease. The distal portion of the peripheral
root was extended under the cylinder wall, so the end of the root was outside the cylinder. We determined that the inside of the cloning cylinder and the dish outside the cylinder were separate compartments by showing that blue dye placed in one compartment did not diffuse into the other compartment over 24 h. In addition, different heights of the liquid medium could be maintained in the two compartments. When control medium was in the distal compartment, Schwann cells migrated with axons that extended from the end of the peripheral nerve (Fig. 3 c). After 24 h, axons and Schwann cells had extended hundreds of micrometers from the nerve root. If 20 μM CHI was in the distal axon compartment, axons still elongated for many hours (Fig. 3 d), but Schwann cells did not leave the nerve roots (Fig. 3, compare c, d). Because of the separation of the compartments, axons and Schwann cells migrated from the explants in the perikaryon compartment of dishes with CHI in the distal axon compartment (Fig. 3 e, f are similar, despite CHI in the axon compartment). Axonal growth rates were comparable in untreated (144 ± 13 μm/h) or CHI-treated (140 ± 14 μm/h) distal axonal compartments. Thus, although axonal growth from DRG explants stops after 1 h of global inhibition of protein synthesis, protein synthesis is not required in distal axons for DRG axonal growth for >24 h.

**Induction of growth cone collapse by repulsive cues does not require protein synthesis**

Growth cone collapse assays were conducted with E7 and E13 DRG explants that were exposed to Sema3A, E8 sympathetic chain ganglia exposed to Sema6A, and E7 temporal retinal explants exposed to negative cues, ephrin-A2 and slit-3 (Table 2, Fig. 4). Explants were cultured 24 h in laminin-coated wells, and then 20 μM CHI or anisomycin or control vehicle were added for 15 min before adding Sema3A to DRG cultures, or ephrin-A2 or slit-3 to temporal retinal cultures for 30 min. As illustrated in Figure 4 and presented in Table 2, the low collapse rates of untreated, 144 ± 13 μm/h) or CHI-treated (140 ± 14 μm/h) distal axonal compartments. Thus, although axonal growth from DRG explants stops after 1 h of global inhibition of protein synthesis, protein synthesis is not required in distal axons for DRG axonal growth for >24 h.

**Figure 2. Nascent protein synthesis in E7 DRG neurons.** The methionine analog AHA in methionine-free DMEM was added to dissociated cells treated with 40 ng/ml NGF (a), 0.5 μg/ml Sema3A (b), or control medium (cont) (c) for 60 min. Cells in d received no AHA. Cells were fixed and incubated with fluorescent alkyne to label the AHA incorporated into nascent proteins. e, Mean staining intensity of NGF- and Sema3A-treated distal axon segments relative to unstimulated neurons. Mean ± SEM. *p < 0.05, **p < 0.001. f–i, The protein synthesis inhibitors CHI and anisomycin (aniso) inhibit protein synthesis. Dissociated DRG cells cultured overnight in medium with 10 ng/ml NGF were incubated 60 min in methionine-free DMEM with AHA (f) or with addition of 20 μM CHI (g) or 20 μM anisomycin (h). Cells in i received no AHA. Fluorescent alkyne labeling of the CHI- and anisomycin-treated neurons and axons (f, g) was as weak as labeling of cells that were not incubated with AHA (h). Scale bars: a (for a–d), 10 μm; f (for f–i), 15 μm. j, Mean staining intensity of distal axons. Mean ± SEM. *p < 0.00001.

**Without protein synthesis inhibitors.** In addition to studies with chick DRGs, we investigated whether protein synthesis was required for growth cones of embryonic mouse DRG neurons to respond to Sema3A. DRGs were explanted from E15 mice and...
cultured 24 h on laminin. Cultures were treated for 15 min with CHI or the vehicle for 15 min, before adding Sema3A. As shown in Figure 4m–p and Table 2, CHI treatment did not inhibit the collapse response of mouse DRG growth cones to Sema3A. Some E7 DRG explants were cultured 8 d before treatment. Many growing axons with active growth cones were present with a mean growth rate of 64 ± 5 μm/h (n = 22), and the growth rate did not change, when 20 μM CHI was added for 30 min (69 ± 4 μm/h, n = 22). When 1 μg/ml Sema3A was added to the CHI-treated dishes, 95% growth cones rapidly collapsed. Thus, protein synthesis is not required for growth cone collapse by several neuronal types in response to several negative cues.

We tested the effects of inhibiting protein synthesis on retinal growth cone responses to surface-bound ephrin-A2. Contact with 6 μM ephrin-A2-coated beads induced 90% of temporal retinal growth cones to collapse or turn away from the beads (n = 20) (Weinl et al. 2005), whereas contacts with BSA-coated beads induced no turning or collapse (n = 8). In the presence of CHI added simultaneously with beads to E7 retinal explants, growth cones did not collapse after contact with BSA-coated beads (n = 9), whereas 95% of CHI-treated growth cones (n = 23) collapsed or turned away after contacting ephrin-A2 beads (Fig. 4q–v). These results, consistent with responses to soluble repulsive cues, indicate that protein synthesis is not required for responses to repulsive guidance cues. The loss of F-actin induced by repulsive cues does not require protein synthesis

Downstream signaling from guidance cue receptors regulates actin filament dynamics via Rho GTPases (Wahl et al., 2000; Hu et al., 2001; Guan and Rao, 2003). Because it was reported that local RhoA synthesis is required for Sema3A-induced growth cone collapse (Wu et al., 2005), we investigated the effects of ephrin-A2 and Sema3A on actin filament organization in control and CHI-treated retinal and DRG growth cones. E7 retinal and DRG explants were cultured on L1, because axonal retraction induced by repulsive cues is diminished on L1, although growth cone collapse still occurs. After 15 min pretreatment with CHI or the control vehicle, explants were exposed to repulsive cues for 15 min, fixed, and stained with rhodamine-phalloidin to label F-actin. Figure 5 shows images and line scans of the distribution of F-actin at the leading margins of control, ephrin-A2, or Sema3A-treated retinal and DRG growth cones, respectively, without and with CHI. Control growth cones had a peak of actin filament density at the leading margin, followed by a flat level of filament density farther back in the growth cone (Fig. 5a,c,e,j). Actin filament distribution and density in growth cones treated 30 min with CHI was identical to controls (Fig. 5b,g). In growth cones treated 15 min with ephrin-A2 (temporal retina) or Sema3A (DRG), the peak in actin filament density at the leading edge was absent, and, as seen previously (Fan et al., 1993), actin filament density was significantly lower throughout the leading margin (Fig. 5c,e,h,j). Growth cones of neurons treated 15 min with CHI and then 15 min with ephrin-A2 or Sema3A and CHI were not different from growth cones treated with repulsive cues alone (Fig. 5d,j). Thus, protein synthesis is not required for signaling downstream from repulsive cues that reduces actin filament content. It is reported that Sema3A stimulates RhoA mRNA translation, and Sema3A increased RhoA density in rat DRG growth cones (Wu et al., 2005). We conducted similar determination of RhoA staining intensity in growth cones of DRG neurons treated with Sema3A with or without inhibiting protein synthesis. As shown in Figure 5k–o, we found no significant changes in mean RhoA staining intensity in DRG growth cones after Sema3A treatment in control or CHI-treated dishes. Thus, treatment of chick DRG neurons with Sema3A did not significantly change the RhoA content of growth cones. The induction of growth cone spreading, increased F-actin, and growth cone turning by NGF does not require protein synthesis

We examined the effects of CHI on responses of E7 DRG growth cones to attractive cues NGF and NT-3. E7 DRG explants were cultured overnight in L1-coated wells in F-12 and B27 without neurotrophins. Growth cone behaviors were recorded before and after adding 40 ng/ml NGF (Fig. 6a) (supplementary Videos S1, S2, available at www.jneurosci.org as supplemental material) or NT-3 (Fig. 6b). Growth cones rapidly expanded in response to either neurotrophin. As measured in Figure 6, c and d (supplementary Videos S3, S4, available at www.jneurosci.org as supplemental material), this expansion response was not inhibited by 15 min pretreatment with CHI, which remained in the media with the neurotrophins. In another experiment, we extended the CHI pretreatment period to 2 h before adding NGF. We observed that, after 2 h in CHI, the spreading response of growth cones to NGF addition was absent (supplemental Fig. S2, available at www.jneurosci.org as supplemental material). Thus, chick embryo growth cones normally contain sufficient components to spread in response to neurotrophins without need for protein synthesis. We next determined the turning response of E7 DRG growth cones toward a soluble NGF gradient. When an NGF-coated micropipette was placed near growth cones, growth cones turned toward the micropipette with a mean turning angle of 12 ± 2.5° (n = 17) (Fig. 6e). Using criteria of Ming et al. (1997) (>5° is a positive turn, −5° to +5° is no turn, less than −5° is a negative turn), 65% of turns were positive, 35% did not turn, and 0% were negative turns. Turning response to a BSA-coated pipette was −0.6 ± 2.4° (n = 9), significantly different from the NGF response (p < 0.005; 11% of turns were positive, 11% negative, and 78% did not turn). We then conducted assays in the presence of CHI, which was added 15 min before placing the micropipette. As illustrated in Figure 6f, growth cone turning toward the pipette occurred with a mean turning angle of 5.4 ± 2.1° (n = 19; 42% positive turns, 53% no turns, and 5% negative turns). This was statistically different from the turning angle of non-CHI-treated growth cones toward NGF (p < 0.05, t test). However, the turning response of CHI-treated growth cones toward a BSA-coated
pipette was -1.7 ± 2.2° (n = 11; 0% positive turns, 91% no turns, and 9% negative turns), which was statistically different from the 5.4 ± 2.1° turning response to an NGF micropipette (p < 0.05). Thus, global inhibition of protein synthesis did reduce growth cone turning toward NGF over a 45 min assay, although the attractive turning response was still present.

Asymmetric distribution of actin filaments is an early phase in growth cone turning toward NGF (Gallo and Letourneau, 2000). We assessed the effects of an NGF gradient on F-actin distribution in live control and CHI-treated growth cones, transfected to express a fluorescent F-actin probe, GFP–UtrCH. This probe contains the F-actin binding domain of utrophin and binds F-actin without altering the dynamics of actin-dependent events (Burkel et al., 2007). Figure 6 shows the average GFP–UtrCH intensity values along 15-pixel-wide line scans taken across the width of growth cones before and after being exposed to BSA, NGF, or NGF gradients in the presence of CHI. A shi in F-actin distribution toward the NGF source was seen in both control and CHI-treated growth cones but not toward a BSA source. A quantification of the F-actin distribution across the growth cones was estimated by summing the average GFP–UtrCH pixel intensities along the line scans in the growth cone halves toward the NGF source and in the halves away from the NGF source and then computing a proximal/distal ratio of the summed F-actin intensities. As a measure of change in F-actin distribution, the proximal/distal intensity ratios of images acquired 2 min after introducing NGF were divided by the ratios determined from images of the same growth cones acquired just before introducing NGF. For control growth cones, this ratio was 1.31 ± 0.15 SE (n = 10), and, for CHI-treated growth cones, the ratio was 1.15 ± 0.09 SE (n = 10). These values were significantly different from ratios computed for growth cones exposed to a local BSA source (0.94 ± 0.06 SE) but were not significantly different from each other by either t test or Mann–Whitney U test. Thus, introduction of a local NGF source induced a shift in growth cone F-actin distribution toward the NGF source, and this was not affected by global inhibition of protein synthesis.

Growth cone spreading is correlated with increased F-actin content in NGF-treated growth cones. To examine changes in actin filament content in DRG growth cones after adding NGF, E7 DRG explants were exposed to global NGF for 15 min and then fixed and stained with rhodamine-phalloidin to quantify F-actin. As shown in Figure 7, the total integrated staining for F-actin in DRG growth cones was increased 250% at 15 min after adding NGF (Fig. 7a,h,i). This increase in F-actin was unaffected by 15 min pretreatment with CHI before and during NGF treatment (Fig. 7c,d,f). Thus, protein synthesis is not required...
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Table 2. Effects of protein synthesis inhibitors on growth cone collapse

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<th># GC counted</th>
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Explanets of E7 chick DRG, temporal retina, sympathetic ganglia, or E15 mouse DRG ganglia were cultured on laminin in F-12 with B27. NGF at 10 ng/ml was added to DRG and sympathetic cultures. After 24 h, explants were treated 15 min with vehicle or 20 μM CHI or anisomycin, followed by 30 min incubation with guidance cues in continued presence of vehicle or protein synthesis inhibitor. Explants were fixed, and growth cones were scored as collapsed, if they had one filopodium or less. The Mann–Whitney U test was used to assess whether protein synthesis inhibitors significantly altered the collapse data of either vehicle-treated or guidance cue-treated cultures. The test indicated that protein synthesis inhibitors had no effect on intrinsic growth cone collapse and only on responses to repulsive guidance cues.

for the rapid increase in F-actin and growth cone spreading that is induced by NGF.

It is reported that 5 min treatment with netrin or BDNF increased the β-actin content of Xenopus growth cones (Leung et al., 2006; Yao et al., 2006). We treated DRG explants with NGF for 15 min with or without 15 min CHI pretreatment and then fixed and stained the cultures with β-actin antibodies (Fig. 7g–j). By quantitative immunofluorescence, we found that NGF treatment did not increase the total integrated staining for β-actin in DRG growth cones and distal axons (Fig. 7k). As noted above, NGF treatment did result in increased F-actin content (Fig. 7f), and β-actin distribution was altered in the spread of NGF-treated growth cones, being concentrated at the leading edge. CHI treatment did not alter total integrated anti-β-actin staining of growth cones, whether exposed or not exposed to NGF (Fig. 7i–k). Thus, NGF treatment did not change the β-actin content of DRG growth cones and distal axons.

**Growth cone responses to guidance cues in compartmented dishes**

We used compartmented cultures to ask whether 24 h inhibition of distal protein synthesis affects growth cone collapse in response to Sema3A. When 1 μg/ml Sema3A was added to axonal compartments treated 24 h with CHI, 73% of growth cones of E7 DRG neurons collapsed within 10 min (n = 24) (Fig. 8a,b). We found that the spreading response of E7 DRG growth cones to global NGF addition also occurred after inhibiting axonal protein synthesis for 24 h (Fig. 8c–f).

To investigate whether protein synthesis in axons is required for a long-term turning response to NGF, we used the compartmented culture and placed agarose beads soaked in either NGF or BSA into a collagen matrix that was formed in the axonal compartment (Fig. 8g,h). The collagen matrix included either 20 μM CHI or the control vehicle. After 24 h, we analyzed axon orientation with respect to the NGF or BSA beads. We drew a line halfway between the beads and measured the orientation angle of the terminal 200 μm of each axon with respect to the closer bead (Fig. 8g,h). In gels with control vehicle, the orientation angle to a BSA bead was 14 ± 7°, whereas orientation to the NGF bead was 46 ± 4°. These values were statistically different (p < 0.0005). In gels containing 20 μM CHI, axon orientation toward a BSA bead was 11 ± 8°, whereas orientation angle toward an NGF bead was 65 ± 7° (p < 0.0001). Thus, DRG axons elongating from a nerve root into collagen could orient toward an NGF bead despite the continuous presence of CHI. These results show that axonal protein synthesis is not required for DRG growth cones to be guided by an NGF gradient.

**Discussion**

Axonal protein synthesis supports multiple functions in adult terminals, including neurotransmission, plasticity, and regeneration (Zheng et al., 2001; Klann and Dever, 2004; Piper and Holt, 2004; Verma et al., 2005; Twiss and van Minnen, 2006; Jiménez-Díaz et al., 2008). Protein synthesis in developing axons totals ~5% of neuronal protein (Eng et al., 1999; Lee and Hollenbeck, 2004). Thus, axonal protein synthesis is not required for DRG growth cones to be guided by an NGF gradient.
However, studies with compartmented cultures indicate that axonal growth does not require axonal protein synthesis (Eng et al., 1999; Blackmore and Letourneau, 2007). Recent papers reported that immediate local protein synthesis is required for responses to guidance cues (Lin and Holt, 2007). We investigated this requirement for chick retinal, DRG, and sympathetic neurons and mouse DRG neurons. Unlike other reports, we found that protein synthesis is not required for chick and mouse DRG and chick retinal and sympathetic growth cones to respond to ephrin-A2, slit-3, semaphorins, NGF, and NT-3. Our assays included cues presented globally and cues locally presented on beads or in a gradient. Furthermore, in novel studies in a compartmented dish in which distal axons were in the continuous presence of CHI but neuronal perikarya were not, axons elongated for hours and growth cones responded to guidance cues in the absence of local protein synthesis. Our results are consistent with two recent publications reporting that protein synthesis is not required for chick retinal growth cone responses to ephrin-A5 (Lang et al., 2008) or for recruitment of DCC (deleted in colorectal cancer) receptors to growth cone surfaces (Bouchard et al., 2008).

Like previous reports, NGF and Sema3A stimulated the phos-
phorylation of the translation repressor eIF-4EBP1. Likewise, nascent protein synthesis in DRG neurons was increased by global addition of NGF or Sema3A. Thus, our findings taken with previous findings (Campbell and Holt, 2001; Takei et al., 2001; Cox et al., 2008) suggest that common mechanisms regulate protein synthesis in developing axons in response to extrinsic factors. However, the relative roles of distally synthesized proteins in axonal functions may depend on metabolic, developmental, or physiologic circumstances of different neurons.

When protein synthesis inhibitors were globally applied to DRG or retinal explants, the axonal growth rate was normal for 1 h, but, eventually, elongation slowed, growth cone motility diminished, and axons retracted. Axonal growth may stop because axonal assembly eventually uses all available components. It may also be that proteins with critical roles in growth cone motility are degraded. We noted that growth cones began to detach and retract during prolonged protein synthesis inhibition. Adhesion receptors that mediate growth cone traction are reutilized via endocytic/exocytic recycling (Caswell and Norman, 2006; Dequidt et al., 2007; Tojima et al., 2007). Perhaps, when protein synthesis is blocked, degradative events during recycling exhaust adhesive receptors. In support of this, growth cone detachment and axonal retraction that occurred after prolonged CHI treatment was delayed, when we also inhibited protein degradation.

When protein synthesis inhibitors were in the axonal compartment of a compartmented chamber, axonal elongation was undiminished for 24 h or more. This indicates that all proteins necessary for axonal growth are made in perikarya and transported distally. As in previous studies (Eng et al., 1999; Blackmore and Letourneau, 2007), axons in our cultures grew several millimeters without distal protein synthesis. Axons in developing embryos reach their targets after similarly growing several millimeters or less. However, adult projection axons can be 10–100 times longer than when they connected to their targets. Even at these lengths, fast axonal transport can deliver vesicle-bound proteins to terminals in a few hours, but proteins that travel via slow transport, especially cytoskeletal components, take days or weeks to reach ends of long axons. Thus, protein synthesis in adult axons is critical to replace degraded proteins, maintain cytoskeletal integrity, mediate plasticity, and support regeneration (Koenig and Giuditta, 1999; Campenot and Eng, 2000; Zhang and Poo, 2002; Piper and Holt, 2004; Verma et al., 2005; Twiss and van Minnen, 2006; Hengst and Jaffrey, 2007).

Why are our results different from reports that local protein synthesis is required for growth cone responses to guidance cues? One possibility involves the neurons and animal species involved. We used DRG neurons, sympathetic neurons, and retinal neurons from embryonic chicks and mice. Other laboratories used embryonic Xenopus retinal (Campbell and Holt, 2001; Leung et al., 2006; Piper et al., 2006) or spinal cord (Guirland et al., 2003; Yao et al., 2006) neurons, and one study used fetal rat DRGs (Wu et al., 2005). Retinal and DRG neurons are common between these studies, so it is unclear whether these differences are attributable to different neuronal phenotypes. The differentiation state of Xenopus spinal cord neurons is uncertain, because the spinal

Figure 5. CHI treatment does not affect Sema3A- and ephrin-A2-induced loss of F-actin, and Sema3A does not increase RhoA content of DRG growth cones. E7 DRG explants were cultured overnight in F-12, B27, NGF (a–d, k–n), and retinal explants were cultured in F-12, B27 (f–l). Sema3A was added to DRG explants for 15 min with (d, n) or without (c, m) 15 min CHI pretreatment. Ephrin-A2 was added to retinal explants for 15 min with (j) or without (h) 15 min CHI pretreatment. Vehicle-treated (a, f, k) and CHI-treated (b, g, l) control explants (Cont) were treated 30 min without repulsive cues. e, f, Mean F-actin staining intensities along lines scanned for 6 μm backward from the growth cone leading edges. Mean ± SEM. g, j, Mean pixel intensity of anti-RhoA staining within traced growth cone borders. Scale bar, 10 μm. Mean ± SEM, no significant differences.

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cords were from young embryos, and the neurons were uncharacterized.

Another source of differences may lie in different levels of metabolism in these in vitro studies. Perhaps, our neurons had high metabolic rates that generated sufficient proteins in the perikarya to supply all growth cone functions. In support of this, after 24 h culture, E7 DRG axons had extended >2000 μm, and E7 retinal axons had extended >1000 μm. E7 DRG explants maintained axonal growth rates of >1 mm/d for at least 8 d. These lengths far exceed axon lengths in published figures of Xenopus retinal and spinal neurons or rat DRG neurons. Xenopus spinal cord neuronal axons were 100–200 μm long and were

Figure 6. Effects of CHI on growth cone spreading and turning in response to NGF and NT-3. E7 DRG explants were cultured overnight in F-12, B27 on L1. Images were acquired immediately before and 10 min after adding 40 ng/ml NGF (a) or NT-3 (b). CHI was added for 15 min before and during treatment with NGF (c) or NT-3 (d). c, d. Mean growth cone area in pixels at 0, 2, and 10 min treatment with NGF, *p < 0.0001, †p < 0.05, ‡p < 0.001 (c) or NT-3, *p < 0.002 (d). Mean ± SEM. e. Time-lapse images of growth cones (arrowheads) turning toward an NGF micropipette. The first image was 10 min before introducing the pipette, and the following images were 2, 10, 20, and 30 min later. f. Time-lapse images of the attractive turning (arrowheads) of growth cones that were treated with CHI 15 min before and during exposure to an NGF gradient from a micropipette. g. GFP–UtrCH distribution in a growth cone before and 2 min after introduction of an NGF micropipette to the left of the growth cone. h. The average intensity of GFP–UtrCH fluorescence intensity along line scans drawn across the width of growth cones before and 2 min after introduction of a micropipette that locally releases BSA or NGF. Growth cones in the farthest right panel were exposed to CHI 15 min before introducing the NGF micropipette. Scale bar: a, b, e–g, 50 μm.
cultured on poly-lysine (Guirland et al., 2003; Yao et al., 2006), unlike the natural substrates we used. Our liquid medium was F-12 with B27 additives, whereas *Xenopus* neurons were cultured in diluted Leibovitz medium with 1% serum and Ringer’s solution. The importance of specific medium components to neuronal metabolism has been appreciated for many years (Bottenstein and Sato, 1979; Chen et al., 2008). Axonal growth by *Xenopus* spinal neurons is most vigorous in the first 12 h, most growth occurs in the first 24 h, and few neurons survive 4 d (Tabi and Poo, 1991). Another indication of high activity in our neurons was the rapidity of growth cone responses to guidance cues. After adding NGF, growth cone spreading and increased F-actin was evident by 2 min, whereas retinal and DRG growth cones began to collapse within 2 min of adding ephrin-A2 or Sema3A. This rapid response is more consistent with mechanisms that activate signaling pathways rather than protein synthesis. *Xenopus* retinal growth cone turning toward netrin was not seen until 10 min after netrin-induced actin synthesis was detected (Leung et al., 2006).

Perhaps, growth cones in our cultures were rich in proteins that function in growth cone motility, whereas levels of these proteins were low in other studies. We found that 45 min CHI treatment produced no significant decreases in growth cone content of RhoA or β-actin, as expected if protein levels are high. Neither did 15–30 min treatment of DRG explants with Sema3A or NGF induce detectable increases in RhoA or β-actin content of growth cones, respectively. Leung et al. (2006) reported that, in *Xenopus* retinal growth cones, β-actin staining intensity was 30% higher after 5 min netrin treatment. Assuming that a retinal growth cone contains 500–1000 ribosomes (Campbell and Holt, 2001) and it takes 1 min to synthesize an actin molecule (Trachsel, 1991), these growth cones would have to initially contain \( \frac{1}{1000} \times 10^7 \) actin monomers for the addition of 5000 monomers (1000 molecules/min for 5 min) to constitute a 30% increase. Actin and tubulin are the most abundant proteins in chick brain growth cones (Cypher and Letourneau, 1991), and, based on cytoplasmic globular actin pools in chick brain neurons (30–37 \( \mu \)M) (Devineni et al., 1999), we estimate that a 10 \( \times \) 20 \( \times \) 1 \( \mu \)m DRG growth cone contains \( \sim 4 \times 10^6 \) β-actin monomers, 100\( \times \) more than *Xenopus* retinal growth cones. In the studies of Wu et al. (2005), rat DRG explants were cultured for 3 d, and the rat
DRG axons had reached 400–500 μm long, which is one-quarter the length of our E7 chick DRG axons after 20 h. As seen in the study of Wu et al. (2005), their Figure 3a, RhoA was barely detectable in embryonic rat DRG growth cones. Conversely, our DRG growth cones stained robustly for RhoA with or without Sema3A treatment, and we found that migrating growth cones in 8 d in vitro DRG cultures collapsed rapidly in response to Sema3A, even with CHI treatment. Our results are consistent with the hypothesis that growth cones of actively growing axons contain sufficient RhoA and β-actin to respond to guidance cues without protein synthesis. Although our data show that guidance cues stimulate protein synthesis in axons of E7 DRG neurons, our data also show that these events do not detectably increase growth cone levels of RhoA or β-actin and are not required for responding to guidance cues.

Global treatment with CHI did affect DRG growth cones. Although globally inhibiting protein synthesis for up to 45 min did not significantly affect growth cone responses, discrete changes FIGURE 8. Growth cone responses to Sema3A and NGF are not inhibited by 24 h CHI treatment in axon compartment. E7 DRG explants were cultured overnight in F-12, B27 in compartmented chambers with 20 μM CHI in the axon compartment. a, b, Growth cones 1 min before (a) and 2 min after (b) adding 1 μg/ml Sema3A. Growth cones collapsed. c–f, Growth cones 0 min before (c–f) and 10 min after (c’–f’) adding 40 ng/ml NGF. Increased growth cone spreading is seen. g and h show montages of axon growth from the end of a CHI-treated peripheral root in a collagen matrix containing an NGF- and a BSA-soaked bead. Axons are traced in black. Axon orientation toward the NGF bead, but not the BSA bead, is seen. Scale bar: a, b, 50 μm; c–f, 20 μm; g, h, 150 μm.
from lacking key proteins may occur. After 2 h CHI treatment, DRG growth cones did not spread in response to NGF. Globally applied CHI reduced the turning of DRG growth cones toward NGF. Turning is more complex than growth cone spreading and may involve exocytosis of vesicle-associated proteins (Tojima et al., 2007) that are transported at fast transport rates of 50–400 mm/d (Squire et al., 2008). Such proteins could be moved from the perikaryon to the tip of a 1000 μm axon in 4–30 min (or 30 s to 3 min in a 100 μm Xenopus spinal neuron axon), which is within the duration of a turning response. Thus, because vesicle transport is rapid, globally applied protein synthesis inhibitors cannot distinguish a growth cone location from a cell body location for protein synthesis that is necessary for growth cone turning. In an attempt to eliminate contributions of perikaryon-derived proteins to growth cone responses, some studies isolated growth cones from cell bodies by severing axons (Campbell and Holt, 2001; Ming et al., 2002; Wu et al., 2005). However, the isolated axons were damaged, because most did not recover growth cones (Ming et al., 2002). In another study of severed axons (Zheng et al., 2001), cycloheximide treatment induced retraction of isolated axons, but axons of intact neurons did not retract after cycloheximide treatment.

Therefore, unlike previous reports, we used compartmented dishes to examine whether distal axonal protein synthesis is required for responses to guidance cues. Because DRG growth cones responded to globally applied Sema3A or NGF and could retract after cycloheximide treatment.

In summary, we confirmed that guidance cues stimulate protein synthesis in distal axons. Protein synthesis may occur in axons from the earliest neuronal polarization and axonal differentiation, via targeted mRNA transport into axons and signaling by extrinsic cues to regulate mRNA translation (Kiebler and Bas- sell, 2006). Extrinsic cues such as Sema3A and NGF may regulate axonal mRNA translation throughout this period, during navigation to targets, later as axons branch, and are remodeled into terminal fields, and in mature organisms to maintain terminals and to respond to stimuli or injury. Our in vitro results indicate that proteins made in distal axons of chick neurons are not necessary for axonal elongation and growth cone guidance cues, whereas other in vitro studies indicate that distally synthesized proteins are required for responses to guidance cues. As noted above, higher metabolic activity in our neurons may explain these differences in in vitro studies. Perhaps, in embryos the relative contributions of locally translated proteins such as RhoA and β-actin to total growth cone pools is initially low as immature axons rapidly elongate to their targets. Later, the significance of distal axon protein synthesis may increase, as axons reach their targets, elongate by intercalary growth to become the greatest part of neuronal mass (Pfister et al., 2004), and form mature endings in a complex environment. Local protein synthesis may become highly important as axon terminals become locally distinct in form and function. In vivo studies across the time span of development are needed to more clearly understand the roles of distal protein synthesis in axonal morphogenesis.

References


