Axonal targeting of trophic receptors is critical for neuronal responses to extracellular developmental cues, yet the underlying trafficking mechanisms remain unclear. Here, we report that tropomyosin-related kinase (Trk) receptors for target-derived neurotrophins are anterogradely trafficked to axons via transcytosis in sympathetic neurons. Using compartmentalized cultures, we show that mature receptors on neuronal soma surfaces are endocytosed and remobilized via Rab11-positive recycling endosomes into axons. Inhibition of dynamin-dependent endocytosis disrupted anterograde transport and localization of TrkA receptors in axons. Anterograde TrkA delivery and exocytosis into axon growth cones is enhanced by nerve growth factor (NGF), acting locally on distal axons. Perturbing endocytic recycling attenuated NGF-dependent signaling and axon growth while enhancing recycling conferred increased neuronal sensitivity to NGF. Our results reveal regulated transcytosis as an unexpected mode of Trk trafficking that serves to rapidly mobilize ready-synthesized receptors to growth cones, thus providing a positive feedback mechanism by which limiting concentrations of target-derived neurotrophins enhance neuronal sensitivity.

Introduction

Polarized membrane trafficking of signaling receptors is critical for developing neurons to respond to extracellular stimuli mediating growth, guidance, and neuronal survival. The family of tropomyosin-related kinases (Trk) receptors provides a prominent example of trophic factor receptors that undergo long-distance axonal trafficking to regulate growth and survival of developing neurons (Zweifel et al., 2005). A key mechanism by which the prototypical neurotrophin, nerve growth factor (NGF), secreted by peripheral target tissues, provides trophic support to sympathetic neurons is by regulating endocytosis and retrograde trafficking of NGF:TrkA complexes in signaling endosomes from axon terminals to cell bodies (Delcroix et al., 2003; Ye et al., 2003; Heerssen et al., 2004; Zweifel et al., 2005). However, the neurotrophin-mediated developmental processes of axon growth, guidance, and neuronal survival must also place an enormous demand on rapid axonal targeting of Trk receptors to ensure sensitive functional responses. Currently, little is known about the membrane trafficking mechanisms regulating proper localization of Trk receptors in axons.

Membrane proteins are often delivered to axons via the secretory pathway (Horton and Ehlers, 2003). Proteins, after synthesis in the rough endoplasmic reticulum (ER) and undergoing maturation in the ER–Golgi network in neuronal cell bodies, are shipped directly to axons in Golgi-derived vesicles. However, in some cases, axonal targeting is accomplished by a more circuitous endocytosis-dependent mechanism called transcytosis in which proteins are initially delivered to somatodendritic compartments and then endocytosed and transported anterogradely for insertion into axon terminals (Horton and Ehlers, 2003). Transcytosis has been best characterized in polarized epithelial cells, including hepatocytes and enterocytes, in which endocytic trafficking from basolateral to apical surfaces is the primary mode for delivery of newly synthesized apical membrane proteins (Bastaki et al., 2002; Tuma and Hubbard, 2003). So far, transcytotic delivery from neuronal soma surfaces to axons has been demonstrated for only a limited number of membrane proteins, including L1/neurona-glial cell adhesion molecule (NgCAM) (Wisco et al., 2003), a cell adhesion molecule important for axon guidance, transferrin (Tf) receptor (Hémar et al., 1997), which regulates cellular iron homeostasis, and the type 1 cannabinoid receptor CB1R (Leterrier et al., 2006), a highly abundant G-protein-coupled receptor in the nervous system implicated in modulating synaptic plasticity. However, the implications of transcytosis on receptor signaling and function have remained unclear.

Here, we show that Trk receptors undergo transcytosis in developing sympathetic neurons, which involves the endocytic removal of biochemically mature receptors from neuronal cell bodies and retargeting them to axons through recycling endosomes. We provide evidence that neurotrophins acting on axon terminals of compartmentalized cultures can recruit their Trk receptors to their sites of action via transcytosis. Perturbing endocytic recycling attenuated NGF-dependent signaling and functional responses, whereas enhancing recycling increased neuronal responsiveness to NGF. Together, our results suggest...
that this previously uncharacterized mode of TrkA trafficking allows neurons to adjust their sensitivity to the limiting concentrations of NGF found in vivo.

Materials and Methods

DNA and adeno viral constructs

Enhanced green fluorescent protein (EGFP)–Rab11a–wild-type (WT) and EGFP–Rab11a–S25N plasmids were a gift from Dr. Michael Ehlers (Duke University, Durham, NC). EGFP–Rab11a–Q70L was generated from EGFP–Rab11a–WT using site-directed mutagenesis (QuickChange; Stratagene). All EGFP–Rab11a constructs were subcloned into psuttle–cytomegalovirus (CMV) vector (Stratagene). The mcherry–Rab11a construct was generated by subcloning mcherry (kindly provided by Dr. Roger Tsien, University of California San Diego, La Jolla, CA) with Rab11a construct into psuttle–CMV vector. The N-terminal FLAG tag was placed on the 5’ end of TrkBTrkA chimeric receptor (provided by Dr. David Ginty, Johns Hopkins University, Baltimore, MD) by PCR. TrkB:A chimeric receptor has amino acid residues 1–423 belonging to the extracellular domain of TrKB and residues 424–809 belonging to the transmembrane and cytoplasmic domains of TrkA. FLAG–TrkB:A was subcloned into pAdTrack–CMV shuttle vector (Stratagene), which also expresses GFP. TrkA–GFP (gift from Dr. David Ginty) and farnesylated EGFP (Clontech) were cloned into psuttle–CMV expression vectors. Replication-incompetent recombinant adenoviral constructs were generated for EGFP–Rab11a constructs and the FLAG–TrkB:A chimeric receptor using the AdEasy adeno viral vector system (Stratagene). Recombinant viral backbones including the genes of interest were transfected into HEK 293 cells (American Type Culture Collection) using Lipofectamine (Invitrogen). High-titer virus stocks were generated using a CsCl gradient. The recombinant adenoviral construct encoding EGFP (Clontech) were cloned into psuttle–CMV expression vectors. Replication-incompetent recombinant adenoviral constructs were generated for EGFP–Rab11a constructs and the FLAG–TrkB:A chimeric receptor using the AdEasy adeno viral vector system (Stratagene). Recombinant viral backbones including the genes of interest were transfected into HEK 293 cells (American Type Culture Collection) using Lipofectamine (Invitrogen). High-titer virus stocks were generated using a CsCl gradient. The recombinant adenoviral construct encoding dominant-negative dynamin-1 (Dyn-1–K44A) was a gift from Dr. Jeffrey E. Pessin (Albert Einstein College of Medicine, Bronx, NY).

Sympathetic neuron culture and transfection

Superior cervical ganglia were dissected from postnatal day 0.5 (P0.5) Sprague Dawley rats and grown in mass or compartmentalized cultures as described previously (Kuruvilla et al., 2004). Cells were maintained in culture with high-glucose DMEM media supplemented with 10% fetal bovine serum (FBS), penicillin/streptomycin (1 U/ml), and NGF (100 ng/ml; Sigma) for 48 h. Neurons were electroporated using Cellax (E. Pessin et al., 2004). EGFP–Rab11a–Q70L was generated for EGFP–Rab11a constructs and the FLAG–TrkB:A chimeric receptor using the AdEasy adeno viral vector system (Stratagene). Recombinant viral backbones including the genes of interest were transfected into HEK 293 cells (American Type Culture Collection) using Lipofectamine (Invitrogen). High-titer virus stocks were generated using a CsCl gradient. The recombinant adenoviral construct encoding dominant-negative dynamin-1 (Dyn-1–K44A) was a gift from Dr. Jeffrey E. Pessin (Albert Einstein College of Medicine, Bronx, NY).

Cell-surface biotinylation assay

Internalization of TrkA receptors

Two days before the biotinylation assay, NGF was withdrawn from mass cultures as described above. Neurons were biotinylated with a reversible membrane-impermeable form of biotin (EZ-Link NHS-S-S-biotin, 1.5 mg/ml in 1× PBS; Pierce Chemical) at 4°C for 25 min (Kuruvilla et al., 2004). Neurons were washed briefly with 1× PBS containing 50 mM glycine (Sigma) to remove remaining unconjugated biotin. Cells were moved to 37°C to promote internalization under the appropriate conditions for 30 min. To block recycling, cells were treated with monensin (10 μM; Sigma) for 30 min during the internalization process. Neurons were returned to 4°C, and the remaining biotinylated surface receptors were stripped of their biotin tag with 50 mM iodoacetamide (Sigma) to quench excess glutathione. Neurons were lysed with 500 μl of radioimmunoprecipitation assay (RIPA) solution (50 mM Tris-HCl, 150 mM NaCl, 1 mM EDTA, 1% NP-40, 0.25% deoxycholate), and supernatants were subjected to precipitation with 50 μl-immobilized neutravidin–agarose beads (Pierce Chemical) and immunoblotting with a TrkA antibody (Millipore Bioscience Research Reagents).

Surface TrkA levels.

Neurons were briefly treated with a 5 min pulse of NGF (100 ng/ml) in the presence or absence of monensin (10 μM) at 37°C and then quickly moved to 4°C. Surface levels of TrkA were assessed by cell-surface biotinylation at 4°C for 30 min, neutraivdin precipitation, and TrkA immunoblotting.

Transcytosis of TrkA receptors in compartmentalized cultures.

Approximately 250,000–300,000 sympathetic neurons were grown per compartmentalized culture chamber for 7–10 d in vitro (DIV) until axonal projections into the side chambers were evident. Neurons were starved of NGF for 2 d, and surface TrkA receptors in cell body compartments were biotinylated with a membrane-impermeable biotin (EZ-Link NHS-S-S-biotin) at 4°C for 25 min. After incubating the cultures at 37°C for 4 h in the presence or absence of NGF (100 ng/ml) added only to axons, lysates were harvested from cell body and axonal compartments separately and subjected to neutraivdin precipitation and TrkA immunoblotting.

Live-cell antibody feeding assay

Trk receptor internalization.

Sympathetic neurons were infected with adenoviral vectors expressing GFP or FLAG–TrkB: A chimeric receptors. Infected neurons were identified by GFP expression. NGF was withdrawn from the cultures without adding the anti-NGF antibody, and cells were incubated with mouse anti-FLAG antibody (M2, 4.2 μg/ml; Sigma) or Fab fragments derived from the FLAG antibody (10 μg/ml; prepared according to the instructions from Pierce Chemical) for 30 min at 4°C in PBS. Excess antibody was washed off, and cells were moved to 37°C for 30 min under various conditions for internalization. Cells were then washed quickly with PBS and immediately fixed with 4% paraformaldehyde (PFA) in PBS for 30 min at room temperature. Cells were then permeabilized with 0.1% Triton X–100/1% BSA/1× PBS, incubated with fluorescently conjugated anti-mouse secondary antibody for 1 h, and then mounted on slides with Antifade Gold (Invitrogen). Images representing 1 μm optical slices were acquired using a Carl Zeiss LSM 510 confocal scanning microscope equipped with argon (458–488 nm) and helium/neon (543–633) lasers. The same confocal acquisition settings were applied to all images taken from a single experiment. Threshold settings for green and red scans were determined, and the integrated fluorescence values for each channel were quantified. For internalization experiments using M2 anti-FLAG antibody, internalization was quantified as the ratio of anti-FLAG immunofluorescence (red) that colocalized with cytoplasmic GFP (green) relative to the total anti-FLAG immunofluorescence. Weighted coefficients of colocalization between the anti-FLAG and GFP fluorescence were determined by LSM Image Examiner program (Carl Zeiss).

Recycling assay.

Sympathetic neurons grown in compartmentalized cultures and expressing FLAG-tagged TrkB: A receptors were incubated with Alexa-647-conjugated Ca2+–sensitive monoclonal anti-FLAG antibody (M1 antibody, 4.2 μg/ml; Sigma) in PBS supplemented with 1% CaCl2 separately in cell body and axonal compartments for 30 min at 4°C. In Figure 1G, “Surface” culture dishes were left at 4°C, whereas “Internalized” and “Recycled” dishes were moved to 37°C for 30 min to allow for internalization. “Internalized” and “Recycled” dishes were then quickly washed three times with ice-cold 1× PBS containing 1 mM EDTA to strip surface-bound FLAG antibodies that had not internalized. Media containing Alexa-546-conjugated anti-mouse secondary antibody was added to all plates, and “Internalized” and “Recycled” dishes were returned to 37°C whereas “Surface” dishes remained at 4°C, for another 30 min. Cultures were quickly washed with ice-cold 1× PBS and immediately fixed with 4% PFA in 1× PBS. Images were acquired using a Carl Zeiss LSM 510 confocal scanning microscope. Optical sections, 1 μm, were taken, and the same confocal acquisition settings were applied to all images taken from a single experiment. In the “Surface” conditions, surface receptors were quantified as the number of pixels double positive for Alexa-647 and Alexa-546 fluorescence per square micrometer in each cell body or axon (for axons, measurements were taken from a stretch of axon equal to 100–250 μm length). In “Internalized” conditions, internalized receptors were calculated as the number of Alexa-647 fluorescent pixels per square micrometer of cell body or axon. Internalized receptors
were then expressed as a percentage of the total surface receptors for 20–30 cells per experiment. In "Recycled" conditions, recycled receptors were calculated as described previously. The percentage of recycled receptors was calculated using the following formula: $(E - Z)/(C - Z) \times 100$, where $E$ is the colocalization coefficient for Alexa-647–FLAG- and Alexa-546-conjugated secondary antibody fluorescence in the "Recycled" condition. $Z$ is the colocalization coefficient for Alexa-647–FLAG and Alexa-546 fluorescence in the "Internalized" condition. $C$ is colocalization coefficient for Alexa-647–FLAG and Alexa-546 fluorescence in "Surface" condition.

Transcytosis of FLAG–TrkB:A receptors in compartmentalized cultures. Sympathetic neurons expressing FLAG–TrkB:A chimeric receptors were grown in compartmentalized culture chambers assembled on collagen-coated glass coverslips for 7–10 DIV until axonal projections were evident in the side compartments. After withdrawing NGF from the culture media, surface chimeric receptors residing in cell body compartments were labeled under live-cell conditions with mouse anti-FLAG antibody (M2, 4.2 μg/ml) or FLAG Fab (10 μg/ml) for indicated time periods at 37°C. Cultures were washed, fixed, permeabilized, and incubated with fluorescently labeled anti-mouse secondary antibody. Chambers were then detached from the coverslips, ensuring that the cells remained attached. Coverslips were mounted on glass slides, and images were acquired with Carl Zeiss LSM 510 confocal microscope. For colocalization experiments with Rab11, neurons were incubated overnight with rabbit anti-Rab11a antibody (1:1000; Zymed), rabbit anti-Rab4 (1:500; ab13252; Abcam), or mouse anti-Rab7 (1:5000; clone Rab7-117; Sigma). For colocalization experiments with Tf, 50 μg/ml Alexa-546-conjugated human Tf (Invitrogen) was added exclusively to the cell body compartments of compartmentalized cultures along with anti-FLAG antibody. Chambers were incubated for 4 h at 37°C, washed several times with PBS, and then fixed and permeabilized. Anti-FLAG was detected as described above.

Time-lapse imaging of axonal transport and total internal reflection fluorescence analyses of Rab11a and TrkA axonal transport. Sympathetic neurons were grown on poly-lysine-coated glass-bottom dishes (MatTek). Cells were electroporated using Cellaxess CX1 system with 200 ng/ml each of mcherry–Rab11a and TrkA–GFP DNA constructs. Neurons were imaged in Leibovitz L-15 media (Invitrogen) supplemented with 0.5% FBS. Axonal transport of mcherry–Rab11a and TrkA–GFP was visualized using 3-I Marianas wide-field live-cell imaging workstation (Intelligent Imaging Innovations) equipped with dual Cascade II 512 EM cameras for simultaneous two-channel acquisition. Images were acquired using 100× objective (1.45 numerical aperture oil-immersion α-plan-fluoar; Carl Zeiss) every 2 s over 200 frames per movie and

Figure 1. Trk receptors undergo constitutive endocytosis and recycling in sympathetic neurons. A, Constitutively internalized TrkA receptors are sequestered intracellularly during blocking recycling with monensin (10 μM). Membrane proteins were subjected to cell-surface biotinylation. Internalized TrkA receptors were detected by surface stripping of biotin, neutral avidin precipitation, and TrkA Western blotting. B, Densitometric quantification of the results shown in A. Results are the mean ± SEM from seven independent experiments. *p < 0.05, t test. C, Monensin-dependent intracellular accumulation of surface TrkA receptors is attenuated in neurons expressing Dyn–K44A. Cell-surface biotinylation assay was performed in neurons infected with adenoviral vectors expressing GFP and Dyn–K44A, in the presence of monensin or NGF. Normalization for protein amounts in infected neurons and is cytoplasmic. Values have been corrected by subtracting any colocalization observed at 4°C. Results are the mean ± SEM; **p < 0.02, ANOVA, followed by a Tukey’s post hoc test.

D, Live antibody feeding assays in neurons expressing FLAG–TrkB:A chimeric receptors show subcellular localization of receptors under the conditions indicated. Scale bar, 5 μm. E, Quantification of internal accumulation of chimeric receptors under the various conditions indicated in D, by assessing the proportion of colocalization of FLAG immunofluorescence with that of GFP (data not shown), which is coexpressed in infected neurons and is cytoplasmic. Values have been corrected by subtracting any colocalization observed at 4°C. Results are the mean ± SEM from five independent experiments. *p < 0.05 relative to No Ligand condition, ANOVA, followed by a Tukey’s post hoc test. F, Schematic of a live-cell ratiometric assay to measure receptor internalization and recycling. G, Spatial analyses of constitutive TrkA transcytosis in neurons. Representative images of surface, constitutive internalized, and recycled chimeric receptors in cell bodies (top row) and axons (bottom row), as assessed by the live-cell ratiometric assay performed independently in cell body and axon compartments in compartmentalized neuronal cultures. Scale bars: Cell Body, 5 μm; Axon, 20 μm. Graphs represent quantification of Trk internalization (I) and recycling (J) from seven independent experiments. Values are means ± SEM; **p = 0.02, t test.
analyzed as described previously (Deinhardt et al., 2006). The distance traveled by double-positive carriers between two consecutive frames, defined as a single movement, was used to determine instantaneous speed. Carriers were analyzed and tracked using NIH ImageJ. Speeds were binned into 0.2 μm/s and plotted using Prism software (GraphPad Software). Kymographs and movies were generated with Slidebook (Intelligent Imaging Innovations).

NGF-induced TrkA–GFP surface recruitment. Total internal reflection fluorescence (TIRF) analysis of Cos-1 cells and sympathetic neurons expressing TrkA–GFP or farnesylated EGFP was performed on the Mari­anas Live Cell Imaging Workstation using a Carl Zeiss TIRF slider and a 100× objective (1.45 numerical aperture, α-plan-fluoro; Carl Zeiss). Images were taken every 2 s for 200 frames, and change in evanescent-field fluorescence (ΔEFEF) was calculated as described previously (Bezerrides et al., 2004) using NIH ImageJ.

BDNF-induced FLAG–TrkBA surface recruitment. Sympathetic neurons were infected with adenovirus expressing FLAG–TrkBA chimeric receptors. Neurons were incubated with the calcium-sensitive mouse anti-FLAG antibody (M1, 4.2 μg/ml; Sigma) for 30 min at 4°C. Cells were washed and moved to 37°C for 30 min to allow for receptor internalization. EDTA (at 1 mM) washes were used to strip remaining surface receptors of anti-FLAG antibody, and cells were then treated with a 5 min pulse of BDNF (100 ng/ml) in culture media containing Alexa-546-conjugated anti-mouse secondary antibody. Cells were quickly washed with cold PBS and fixed with 4% PFA. Images were taken using LSM 510 confocal microscope.

Axon growth and neuronal survival
Sympathetic neurons grown in compartmentalized culture chambers were infected with adenoviral constructs expressing GFP, EGFP–Rab11a–S25N, and EGFP–Rab11a–Q70L. Axon growth (micrometers per day) in response to NGF added at the indicated concentrations only to axon compartments was measured as described previously (Kuruvilla et al., 2004). To assess neuronal survival, sympathetic neurons from P0.5 rats grown on collagen-coated coverslips for 12 h were infected with adenovirus expressing FLAG–TrkBA receptors. Neurons were grown in the presence of 10 ng/ml NGF for another 48 h. Neurons were then washed and cultured in the presence of BDNF (25 ng/ml) and/or anti-FIAG antibody (M2, 4.2 μg/ml) for 72 h with the addition of anti-NGF (Sigma). Cells were fixed, permeabilized, and incubated with Hoechst 33342 (1:1000) in PBS for 1 h. Neurons with pyknotic, condensed, or fragmented nuclei were scored as dead as described previously (Kuruvilla et al., 2004). Percentage survival is expressed relative to 10 ng/ml NGF (100%).

Immunoprecipitation, immunoblotting, and antibodies
Neurons were lysed with RIPA solution, and lysates were subjected to immunoprecipitation with anti-phosphotyrosine (P-Tyr) (PY-20; Sigma), anti-FLAG (M2; Sigma), or TrkB (BD Biosciences) and incubated with Protein-A agarose beads (Santa Cruz Biotechnology). Immunoblotting analyses of sympathetic neuron lysates were performed as described previously (Kuruvilla et al., 2004). The following antibodies were used for immunoblotting: TrkA (Millipore Bioscience Research Reagents), P-Tyr (PY20; Sigma), and P-Akt (Ser473) and P-Erk1/2 (Thr202/Tyr204) (Cell Signaling Technology). Normalization for protein amounts was done by stripping immunoblots and reprobing for the p85 subunit of phosphatidylinositol 3-kinase (Upstate Biotechnology). Blots were visualized with ECL Plus Detection Reagent (GE Healthcare) and scanned with a Typhoon 9410 Variable Mode Imager (GE Healthcare). Densitometry analysis was performed with ImageQuant software (Molecular Dynamics).

Statistical analyses
Prism software was used for statistical analyses, and all data passed normality tests. All Student’s t tests were performed assuming Gaussian distribution, two-tailed, unpaired, a confidence interval of 95%. One-way or two-way ANOVA analyses were performed when more than two groups were compared.

Results
Trk receptors undergo constitutive endocytosis and recycling in developing sympathetic neurons
Previous studies had indicated that Trk receptors undergo endocytosis predominantly in a ligand-dependent manner (Grimes et al., 1996; Kuruvilla et al., 2004; Chen et al., 2005). We found that TrkA receptors undergo constitutive cycling between the plasma membrane and intracellular endosomes in sympathetic neurons. In a cell-surface biotinylation assay to follow the fate of surface TrkA receptors, NGF stimulation leads to robust internalization of TrkA receptors (Fig. 1A, B), as reported previously. Interestingly, blocking endocytic recycling with the recycling inhibitor monensin (Busu et al., 1981) resulted in significant intracellular accumulation of Trk receptors in the absence of any ligand (Fig. 1A, B), suggesting that surface Trk receptors undergo constitutive internalization and recycling back to the plasma membrane. Blocking recycling in the presence of NGF did not induce any additional intracellular accumulation of TrkA receptors beyond that elicited by NGF alone (Fig. 1A, B). Monensin-dependent intracellular accumulation of TrkA is attenuated in neurons expressing a dominant-negative form (Dyn–K44A) of the GTPase dynamin (Ye et al., 2003), which functions in vesicle scission during endocytosis (Fig. 1C), suggesting that constitutive trafficking of TrkA receptors uses components of the cellular endocytic machinery.

To visualize Trk trafficking in neurons, we also performed live-cell antibody feeding assays in neurons expressing FLAG-tagged chimeric receptors that have the extracellular domain of TrkB and the transmembrane and intracellular domains of TrkA (FLAG–TrkBA). Sympathetic neurons do not normally express TrkB (Atwal et al., 2000), but neurons expressing chimeric receptors showed signaling responses to the TrkB ligand, BDNF, in a manner similar to endogenous TrkA receptors stimulated with NGF (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). The chimeric receptors showed no cross-activation with endogenous TrkA receptors (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Live-cell immunocytochemistry was performed using an antibody directed against the extracellular FLAG epitope of chimeric Trk receptors. We did not observe any signaling or survival responses in neurons treated with antibody alone (supplemental Fig. 1, available at www.jneurosci.org as supplemental material), indicating that the antibodies were not autoactivating. Live-cell immunocytochemistry revealed exclusively surface localization of the receptors at 4°C, a temperature at which little protein trafficking occurs (Fig. 1D). However, at 37°C, the permissive temperature for trafficking, we observed a punctate intracellular localization of Trk receptors, which could be enhanced by either blocking recycling or adding ligand (Fig. 1D,E). Because the FLAG IgG antibodies bind to the extracellular region of FLAG–TrkBA receptors and could potentially induce receptor dimerization and subsequent internalization, we also used, in uptake assays, FLAG Fab fragments that retain the ability to bind antigen without the potential for nonspecific receptor clustering. FLAG-tagged receptors labeled with Fab fragments showed similar internalization and trafficking as those labeled with IgG (supplemental Fig. 2, available at www.jneurosci.org as supplemental material). Together with our findings that the FLAG antibodies by themselves do not induce any signaling or survival responses, these results indicate that internalization observed with live antibody-feeding assays are not an artifact of antibody-induced receptor clustering. Control GFP-expressing sympathetic neurons showed little to no
FLAG-immunoreactivity with either the IgG or Fab fragments (supplemental Fig. 2, available at www.jneurosci.org as supplemental material), pointing to the specificity of uptake of the FLAG antibody in FLAG–TrkB:A-expressing neurons. Together, these results demonstrate a previously unappreciated mode of ligand-independent trafficking of Trk receptors, in which receptors are constitutively endocytosed and recycled in sympathetic neurons.

**Constitutive endocytosis of Trk receptors occurs predominantly in cell bodies**

Given the polarized morphology of neurons, we asked whether constitutive trafficking of Trk receptors differs between cell bodies and axons. Neurons were grown in compartmentalized cultures and a modified live-cell antibody feeding assay (Chen et al., 2005) was used to directly monitor internalization and recycling of cell-surface FLAG–TrkB:A receptors labeled with Alexa-647–FLAG antibodies independently in cell bodies and axons (Fig. 1F). After an incubation period of 30 min, which allowed for the internalization of labeled receptors, surface-bound antibodies were stripped, leaving antibodies bound only to the internalized pool of receptors (Fig. 1G). Additional incubation of cell body and axon compartments for an additional 30 min with Alexa-546 secondary antibodies tagged receptors that recycled to the cell surface (Fig. 1G). Quantification by ratiometric analyses showed that the percentage of surface TrkA receptors that undergo ligand-independent internalization is significantly higher in neuronal cell bodies (59.31 ± 14.7%) than in axons (14.05 ± 6.73%) (\(p = 0.02, t\) test; \(n = 7\) independent experiments) (Fig. 1H). However, of the receptors that internalize, recycling occurs to a similar extent in cell bodies (24.68 ± 5.18%) and axonal (34.18 ± 5.70%) compartments (\(p = 0.24, t\) test; \(n = 7\) independent experiments) (Fig. 1I). These results indicate that ligand-independent internalization of Trk receptors occurs predominantly in neuronal cell bodies but is minimal in axons.

**Neurotrophins acting on axons promote anterograde transcytosis of Trk receptors**

We hypothesized that constitutive endocytosis of Trk receptors in neuronal cell bodies would maintain a dynamic intracellular receptor pool that could be mobilized to axons during neurotrophin stimulation. This would be a way to transport Trk receptors to axons to modulate responses to target-derived neurotrophins without the need for de novo synthesis. To test this hypothesis, we used live-cell antibody feeding in compartmentalized cultures to selectively label and follow Trk receptors originating from surfaces of cell bodies. Neurons expressing FLAG–TrkB:A receptors were grown in compartmentalized cultures, and surface receptors were labeled with anti-FLAG antibody exclusively in cell body compartments (Fig. 2A). Labeling with a fluorescent secondary antibody revealed the presence of FLAG-antibody–bound receptors in axons after a 4 h, but not 30 min, incubation period (Fig. 2B). Because antibody-tagged receptors could only have originated from surfaces of neuronal cell bodies, these results provide direct evidence for anterograde transcytosis of Trk receptors in sympathetic neurons.

To test whether target-derived neurotrophins regulate transcytosis of Trk receptors, we assessed the effects of BDNF added exclusively to axon compartments of compartmentalized cultures on anterograde trafficking of Trk receptors originating from neuronal soma surfaces. BDNF treatment of axons markedly enhanced the levels of anti-FLAG-labeled chimeric Trk receptors in axons after 4 h, with a 5.93 ± 1.14-fold increase compared with unstimulated neurons (Fig. 2B) (\(p = 0.04, t\) test; \(n = 3\) independent experiments). Similar results were obtained when neuronal cell bodies were labeled with anti-FLAG Fab fragments (supplemental Fig. 2, available at www.jneurosci.org as supplemental material).

To determine whether endogenous TrkA receptors in sympathetic neurons undergo a similar process of internalization from neuronal cell bodies and anterograde transport, we biotinylated membrane proteins selectively in cell body compartments of compartmentalized cultures. Neutravidin precipitation followed by Western blot analysis of cell lysates from the cell bodies and axonal compartments showed the presence of biotinylated TrkA receptors in axons after a 4 h incubation period (Fig. 2C), demonstrating that endogenous TrkA receptors in sympathetic neurons undergo transcytosis. The TrkA receptor initially synthesized in the rough endoplasmic reticulum is in a precursor 110 kDa form that, in the course of reaching the plasma membrane, acquires a 30 kDa sugar moiety to become the mature glycosylated form (Jullien et al., 2003). Biotinylated receptors on SDS-PAGE gels corresponded to the ~140 kDa form of the TrkA receptors, indicating that it is the biochemically mature form of the receptor that is available on neuronal cell bodies for surface labeling with biotin and subsequently endocytosed and anterogradely transported to axons. Treatment of axon terminals with NGF enhanced axonal transcytosis of TrkA receptors by 1.32 ± 0.07-fold compared with unstimulated conditions (Fig. 2C) (\(p = 0.03, t\) test; \(n = 4\) independent experiments). The difference between the levels of BDNF-induced axonal transcytosis of FLAG–TrkB:A receptors (5.93 ± 1.14-fold increase) versus NGF-induced transport of endogenous TrkA receptors (1.32 ± 0.07-fold increase) might be explained by that a more sensitive microscopy-based assay was used to detect transcytosing receptors in the first case compared with Western blotting of biotinylated receptors in the second. In addition, for the fluorescence-based assay, neurons were grown in Camp10 Teflon dividers in which the distance between neuronal cell bodies and distal axonal processes is ~1 mm. However, for the biotinylation-based assay, neurons were cultured in larger biochemistry chambers in which the distance between cell bodies and distal processes is ~3–5 mm. Given that, in both cases, appearance of anterogradely transported Trk receptors was assessed after 4 h, more Trk receptors could have accumulated over time in the shorter axons in the Camp10 chambers compared with biochemistry chambers.

Axonal transport of biotinylated TrkA receptors from cell body to axonal compartments is inhibited by expression of Dyn–K44A (Fig. 2D) (60 ± 0.13% decrease compared with control GFP-expressing neurons; \(p = 0.04, t\) test; \(n = 3\) independent experiments), indicating that anterograde movement of Trk receptor is endocytosis dependent and is not attributable to lateral diffusion of receptors along the plasma membrane. To determine whether this mode of trafficking contributes significantly to axonal targeting of TrkA receptors, we examined the effect of Dyn–K44A on axonal levels of TrkA receptors. Blockade of endocytosis reduced axonal levels of TrkA receptors by 43 ± 0.09% (Fig. 2E) (\(p = 0.04, t\) test; \(n = 3\) independent experiments), without affecting total TrkA expression in neurons (Fig. 2F) or surface levels of TrkA receptors on neuronal cell bodies (supplemental Fig. 2, available at www.jneurosci.org as supplemental material). Together, these results suggest that target-derived neurotrophins acting on axon terminals recruit a considerable proportion of Trk receptors to their sites of action by anterograde transcytosis.
immunopositive structures colocalized with endogenous Rab11 in cell bodies and axons, respectively. Transcytosing Trk receptors were also found to colocalize with Tf, a well established marker for the recycling pathway (Fig. 3B). Neurons expressing FLAG–TrkB:A receptors grown in compartmentalized cultures were loaded with anti-FLAG antibodies and Alexa-546–Tf exclusively in cell body compartments. After 4 h of labeling, fixing and permeabilizing of neurons, and staining with a fluorescent secondary antibody for FLAG immunoreactivity, 47 ± 4.5 and 46 ± 5% of FLAG-immunopositive structures colocalized with Alexa-546–Tf in cell bodies and axons, respectively. The degree of colocalization of FLAG–TrkB:A with Rab11 and transferrin was well above that observed with Rab4, a Rab GTPase implicated in a rapid recycling pathway (Maxfield and McGraw, 2004), and Rab7, a Rab protein associated with late endosomes and shown previously to be required for retrograde transport of neurotrophin receptors (Deinhardt et al., 2006) (supplemental Fig. 3, available at www.jneurosci.org as supplemental material).

To further investigate the role of Rab11 in TrkA trafficking, we also performed live imaging of sympathetic neurons expressing TrkA–GFP and mcherry–Rab11. A total of 62 ± 6% of TrkA–GFP puncta were observed to be cotransported with Rab11-positive structures in axons. Time-lapse imaging revealed punctae positive for both TrkA and Rab11 exhibiting highly dynamic bidirectional movements in axons, with an anterograde bias (Fig. 3C) (supplemental Movie 1, available at www.jneurosci.org as supplemental material). A representative kymograph shows punctae positive for both TrkA–GFP and mcherry–Rab11 exhibiting saltatory movement, switching from being stationary to rapid anterograde movement, followed by pausing (Fig. 3C). Kinetic analysis showed that, of the total number of double-positive carriers, ~54% were stationary or pausing. Of the mobile (>0.6 μm/s) double-positive carriers, 66% moved anterogradely, whereas 33% moved retrogradely, with an average speed of 1.2 μm/s in either direction (Fig. 3D).

To specifically address the requirement of Rab11 in axonal transcytosis of Trk receptors, we assessed anterograde transport of Trk receptors originating from neuronal soma surfaces in neurons grown in compartmentalized cultures and infected with adenoviral vectors expressing GFP or a dominant-negative GDP-bound mutant form of Rab11a (Rab11a–S25N) (Park et al., 2004). TrkA transcytosis was assessed by biotinylation of membrane proteins in cell body compartments and detection of transcytosed Trk receptors in axons by neutravidin precipitation and Western blotting. Indeed, expression of Rab11a–

**Rab11-positive recycling endosomes mediate TrkA transcytosis**

Cargo undergoing transcytosis in polarized epithelial cells are commonly transported to their final destinations via recycling endosomes (Altschuler et al., 2003; Tuma and Hubbard, 2003). FLAG antibody labeling of surface Trk receptors in neuronal cell bodies of compartmentalized cultures followed by immunostaining for the recycling endosome-associated small GTPase Rab11a (Zerial and McBride, 2001) showed that transcytosing Trk receptors colocalized or were in close apposition to Rab11-immunoreactive vesicles (Fig. 3A) (50 ± 6 and 45 ± 5% of FLAG-

**Figure 2.** Trk receptors are transcytosed from neuronal soma surfaces to axons. **A,** Schematic of antibody feeding assay in compartmentalized cultures expressing FLAG–TrkB:A to follow transcytosis of Trk receptors originating from neuronal soma surfaces. **B,** Representative images of cell bodies and axonal projections fixed and stained for anti-FLAG antibody. Anti-FLAG-labeled TrkB:A receptors are evident in cell bodies at 30 min and 4 hr (small arrowheads) and in axons only in the 4 hr incubations (large arrowheads). Scale bars, 5 μm. **C,** NGF treatment of axons (100 ng/ml, 4 hr) enhances levels of biotinylated TrkA receptors in axons. Lysates from cell body (CB) and axon (AX) compartments in compartmentalized cultures were precipitated with neutravidin and immunoblotted with TrkA. Normalization for protein amounts was performed by immunoblotting for p85 subunit of PI3K. **D,** Dominant-negative dynamin (Dyn–K44A) inhibits TrkA transcytosis. Cell-surface biotinylation assay was performed in compartmentalized cultures expressing GFP or Dyn–K44A and treated with NGF on axons (100 ng/ml, 4 hr). **E, F,** Dyn–K44A leads to a decrease in axonal, but not total, TrkA levels. Western blot analysis of TrkA levels in compartmentalized cultures (E) shows that expression of Dyn–K44A leads to a significant decrease in axonal TrkA levels. Dyn–K44A does not affect total TrkA levels in neurons (F).
S25N caused a significant reduction (63 ± 0.09% decrease) in levels of biotinylated Trk receptors detected in axons, in the presence of NGF (Fig. 3E) (p = 0.02, t test; n = 3 independent experiments). No significant differences in total levels of TrkA were detected between neurons expressing GFP and Rab11a–S25N, indicating that Rab11 functions in trafficking rather than expression of neuronal Trk receptors (supplemental Fig. 3, available at www.jneurosci.org as supplemental material). Together, these results provide evidence that Rab11-containing recycling endosomes mediate axonal transcytosis of Trk receptors.

NGF stimulation promotes rapid exocytosis of TrkA receptors

We then asked whether neurotrophins, in addition to regulating Trk receptor mobilization to axons, also promote their insertion into the plasma membrane. Previous studies had suggested that TrkB receptors are rapidly recruited to the cell surface in response to neural activity or BDNF (Meyer-Franke et al., 1998; Du et al., 2000; Haapasalo et al., 2002). To determine whether NGF promotes TrkA exocytosis, sympathetic neurons were stimulated briefly (5 min pulse) with NGF, and surface levels of TrkA were
assessed by cell-surface biotinylation. We found that this short stimulus resulted in increased surface TrkA levels, which was attenuated in the presence of the recycling blocker monensin (Fig. 4A, B). Monensin treatment alone, in the absence of ligand, did not appreciably alter surface levels of TrkA within 5 min (Fig. 4B). TIRF analyses also revealed rapid recruitment to the surface of TrkA receptors during NGF stimulation in Cos-1 cells (supplemental Fig. 4, available at www.jneurosci.org as supplemental material), and axonal growth cones of neurons expressing TrkA–GFP (Fig. 4C) (supplemental Movie 2, available at www.jneurosci.org as supplemental material). NGF stimulation induced a significant increase in TrkA–GFP fluorescence at or near the surface (Fig. 4C, D). NGF stimulation did not elicit any changes in surface fluorescence of a membrane-bound marker, farnesylated EGFP (Fig. 4E), suggesting that the observed increase in TrkA–GFP fluorescence is attributable to movement of vesicles relative to the plasma membrane and not attributable to generalized movement or expansion of the plasma membrane.

TrkA–GFP does not distinguish between newly synthesized receptors versus those that once resided on the plasma membrane. Additionally, our TIRF analysis showing increased fluorescence with neurotrophin stimulation at the axonal growth cones does not allow us to determine whether TrkA–GFP receptors are indeed being inserted into the plasma membrane as opposed to receptors merely approaching the plasma membrane. To investigate these two issues, we determined whether receptors being recruited to axonal growth cones with the very brief neurotrophin stimulation originate from a pool that once resided on the plasma membrane and whether they are indeed appearing on the axonal surface. Neurons expressing FLAG–TrkB:A receptors were live labeled with FLAG antibodies and stripped of surface antibodies after allowing internalization at 37°C in the absence of any ligand, and then axonal growth cones were stimulated with the ligand BDNF for 5 min, in the presence of a fluorescent secondary antibody. BDNF treatment significantly increased FLAG immunofluorescence in axon terminals compared with unstimulated neurons (Fig. 4F). Because neurons were treated with the fluorescent secondary antibody under nonpermeabilizing conditions, these results are indicative of delivery to the axonal plasma membrane of recep-

Figure 4. NGF stimulation promotes rapid exocytosis of TrkA from recycling endosomes. A, TrkA is rapidly recruited to the cell surface during NGF stimulation. Levels of surface TrkA were assessed using cell-surface biotinylation in neuronal cultures briefly stimulated with a 5 min pulse of NGF (100 ng/ml) in the presence or absence of the recycling blocker monensin. Normalization for protein amounts was performed by immunoblotting neuronal lysates for p85 subunit of PI3K. B, Densitometric quantification of the results shown in A. Results are quantified relative to the NGF (100 ng/ml) condition, from eight independent experiments. *p < 0.05 and **p < 0.01 as determined by one-way ANOVA, followed by a Dunnett’s multiple comparison test. C, TIRF analyses of growth cones from neurons expressing TrkA–GFP. Dashed lines outline the periphery of growth cones. Scale bar, 5 μm. D, Time course of TrkA–GFP fluorescence in the two growth cones depicted in C. Arrow indicates time of NGF addition. E, Group data showing maximum change in fluorescence (1–5 min) from growth cones of neurons expressing TrkA–GFP (n = 15) or farnesylated EGFP. Values are means ± SEM from four independent experiments. **p = 0.003, t test. F, Rapid BDNF-mediated surface delivery of FLAG–TrkB:A receptors originating from the plasma membrane. FLAG–TrkB:A receptors were live labeled with FLAG antibodies and allowed to internalize, surface antibodies were stripped off, and neurons were stimulated with BDNF for 5 min in the presence of a fluorescent secondary antibody.
tors that had been previously on the cell surface during live labeling with FLAG antibodies. Given that constitutive internalization occurs predominantly in neuronal soma and is minimal in axons (Fig. 1H), we reason that the FLAG-antibody-bound receptors being recruited to axonal membranes originate primarily from soma surfaces. Together, these results suggest that Trk receptors held in intracellular endocytic compartments can indeed be rapidly mobilized to the cell surface during neurotrophin stimulation.

Endocytic recycling modulates neuronal sensitivity to NGF-dependent signaling and axon growth

To examine whether rapid mobilization of Trk receptors to neuronal surfaces via endocytic recycling imparts neuronal sensitivity to NGF, we assessed NGF-dependent signaling in neurons exposed to increasing concentrations of NGF for 5 min, in the presence or absence of monensin. We found that, in neurons treated with monensin, higher concentrations of NGF were required to elicit similar levels of phosphorylation of TrkA and the downstream signaling effectors Akt and Erk1/2 seen in control neurons (Fig. 5A, B). The decrease in P-TrkA levels at high concentrations of NGF (1000 ng/ml) probably reflects rapid degradation of the receptor (Bogenmann et al., 1998), negative regulation of Trk activation by engagement of p75 receptors (MacPhee and Barker, 1997), or recruitment of a phosphatase (Marsh et al., 2003). Together, our results indicate that blockade of receptor recycling attenuates neuronal sensitivity to NGF.

To investigate the role of recycling in functional responses to target-derived NGF, we assessed the effects of a constitutively active GTP-bound variant of Rab11 (EGFP–Rab11a–Q70L) (Uhlig et al., 2006) and the dominant-negative EGFP–Rab11a–S25N mutant on axon growth in compartmentalized cultures when NGF is applied only to axons. Quantification of axon extension (micrometers per day) in response to axon-applied NGF revealed that enhanced recycling with Rab11a–Q70L conferred neuronal sensitivity to lower concentrations of NGF compared with control cultures expressing GFP (Fig. 5C, D). Conversely, neurons exhibit decreased responsiveness to NGF when recycling is impaired by expression of Rab11a–S25N (Fig. 5C, D). Increasing NGF concentration beyond saturating levels eliminated all differences in axon growth between cultures expressing the Rab11 variants and control cultures (Fig. 5D).

Figure 5. NGF-induced TrkA exocytosis modulates neuronal sensitivity to NGF-dependent signaling and axon growth. A, Blocking endocytic recycling attenuates NGF-dependent signaling. Neuronal cultures were briefly stimulated (5 min) with indicated concentrations of NGF in the presence or absence of the recycling blocker monensin (10 μM). Lysates were immunoblotted to detect P-TrkA, P-Erk1/2, and P-Akt. Normalization for protein amounts was performed by stripping the immunoblots and reprobing for p85. B, Densitometric quantification of the results shown in A. Results are quantified relative to the NGF (1000 ng/ml) condition, from seven independent experiments. *p < 0.05 as determined by two-way ANOVA, followed by Bonferroni’s post hoc test. C, D, Rab11a activity regulates NGF-dependent axon growth. Compartmentalized cultures were infected with adenoviruses expressing GFP or Rab11 constructs, EGFP–Rab11a–S25N (dominant-negative) and EGFP–Rab11a–Q70L (constitutively active), and maintained with NGF added solely to the axonal compartments. Rate of axon extension (micrometers per day) was assessed for 72 h. Panels in C are representative images of axons immunostained with anti-β-III tubulin. Separation between cell body (CB) and axon (AX) compartments is indicated by dashed line. D, Graph shows quantification of axon growth. Values are means ± SEM from seven independent experiments. *p < 0.05 and **p < 0.01, two-way ANOVA analysis with Bonferroni’s post hoc test.
target-derived neurotrophins found in vivo amplify neuronal responsiveness to the limiting amounts of NGF. Our findings suggest that regulating this transcytotic trafficking pathway is a mechanism to enhance neuronal responsiveness to NGF. Endocytic recycling attenuated NGF-dependent signaling and by regulating this transcytotic trafficking pathway. Perturbing endocytic recycling facilitated sensitive neuronal responses to the limiting concentrations of target-derived NGF found in vivo.

Discussion

By using compartmentalized cultures in this study, we investigated axonal targeting of Trk receptors in response to target-derived neurotrophins in developing sympathetic neurons. Trk receptors are constitutively endocytosed preferentially in cell bodies, thereby generating a dynamic reservoir of intracellular receptors that can then be delivered to axons via Rab11-containing recycling endosomes. We show that ligand acting locally in axons can recruit receptors from neuronal soma surfaces by regulating this transcytotic trafficking pathway. Perturbing endocytic recycling attenuated NGF-dependent signaling and functional responses, whereas enhancing recycling increased neuronal responsiveness to NGF. Our findings suggest that regulated anterograde trafficking of Trk receptors is a mechanism to amplify neuronal responsiveness to the limiting amounts of target-derived neurotrophins found in vivo.

Currently, the trafficking mechanisms by which signaling receptors are targeted to axons are poorly understood, despite the implications for how neurons might respond to extracellular cues promoting migration, axon growth, guidance, and neuronal survival. Previous studies have shown that, like the targeting of apical membrane proteins in polarized epithelial cells via transcytosis, neurons use endocytic mechanisms for targeting of some axonal proteins. A well-characterized example is L1/NgCAM, a cell adhesion molecule important for axon guidance; initial delivery of L1/NgCAM to the somatodendritic compartment in hippocampal neurons is followed by endocytosis and transport to axonal surfaces via recycling endosomes (Wisco et al., 2003; Winckler, 2004; Anderson et al., 2005; Yap et al., 2008a,b). However, the physiological relevance of such a trafficking pathway in developing neurons and its regulation by extracellular signals still remains elusive.

Based on our findings, we propose that transcytosis might be a more general mechanism than currently appreciated for axonal delivery of receptors in response to extracellular signals impinging locally on axonal growth cones. Thus, target-derived neurotrophins, acting on terminals of axons that can be millimeters long, recruit their own receptors to their sites of action through transcytosis, and this process influences developmental outcomes such as axon growth. Transcytotic trafficking provides the advantage of using ready-synthesized receptors to axons. Membrane trafficking through the endocytic pathway has been reported to be 10 times greater than through the secretory pathway (Horton and Ehlers, 2003), suggesting that the former might provide a larger intracellular reservoir of membrane proteins for regulated delivery to the cell surface. The considerable heterogeneity of endocytic organelles compared with Golgi-derived vesicles might also allow for more plasticity in the regulation of axonal trafficking. We found that Rab11, a small Rab GTPase predominantly found in recycling endosomes, regulates anterograde Trk transcytosis and NGF-dependent axonal extension. Interestingly, protrudin, a Rab11-binding protein, has been implicated in directional membrane trafficking leading to NGF-mediated neurite outgrowth in PC12 cells (Shirane and Nakayama, 2006). However, unlike our results, protrudin requires GDP-bound Rab11 to mediate neurite outgrowth, although we find that axon extension is inhibited by the GDP-bound mutant form of Rab11a (Rab11a–S25N) and enhanced by the GTP-bound Rab11a–Q70L. These results may present an interesting difference between mechanisms that initiate neurite outgrowth from those involved in axonal extension. Although our data indicates that Rab11-positive recycling endosomes mediate axonal transport of Trk receptors, it is likely that receptors internalizing from neuronal soma surfaces have to traverse through several intermediate compartments, including early endosomes and/or multivesicular bodies, before entering axons, as seen for L1/NgCAM (Yap et al., 2008b).

In previous studies of axonal targeting of NgCAM, a point of contention has been whether the protein indeed resides transiently on somatodendritic surfaces, before axonal delivery (Horton and Ehlers, 2003; Sampo et al., 2003; Wisco et al., 2003). By performing cell-surface biotinylation and antibody-feeding assays in compartmentalized cultures, we have provided direct evidence that Trk receptors residing on neuronal soma surfaces are indeed transported anterogradely to axons. These results imply that newly synthesized Trk receptors are first inserted into the plasma membrane of neuronal cell bodies, before being retrieved by endocytosis and anterogradely transported. Insertion of Trk receptors into the plasma membrane of cell bodies before the long journey down the axons might be required to “unmask” an axonal targeting signal, perhaps, via posttranslational modifications or proteolytic processing. In this regard, NgCAM with a point mutation in a tyrosine-based YRSLE motif was found to avoid the transcytotic route and was shipped directly to axons (Wisco et al., 2003), pointing to a role for phosphorylation in regulating transcytosis. It is also tempting to speculate that internalization of unliganded TrkA receptors in neuronal cell bodies acts as a quality control step for the axonal targeting of an “endocytosis-competent” pool of receptors that can then be quickly internalized in response to NGF at the axon terminals. Although retrograde transport of TrkA receptors to mediate NGF-dependent trophic responses has been extensively studied, other modes of Trk trafficking in neurons are poorly understood.
Previous studies examining TrkA recycling have yielded differing results; Chen et al. (2005) showed significant ligand-dependent recycling of TrkA receptors, whereas Saxena et al. (2005) reported very little TrkA recycling when compared with p75 neurotrophin receptors that recycle robustly in response to NGF (Bronfman et al., 2003; Saxena et al., 2005). These studies performed in nonpolarized PC12 cells may not completely recapitulate spatial aspects of receptor trafficking in morphologically complex neurons. We show in sympathetic neurons that Trk receptors undergo ligand-independent internalization primarily in cell bodies, although recycling of internalized receptors is equivalent in cell bodies and axons. Similar to our observations with Trk receptors in sympathetic neurons, p75 receptors recycle in a ligand-independent manner in both cell bodies and axons of motor neurons (Deinhardt et al., 2007), although it remains to be determined whether these receptors are also targeted to axon terminals via regulated transcytosis, as we observed for TrkA. It must be noted here that, although our study highlights a role for long-distance anterograde recycling of unliganded Trk receptors in amplifying neurotrophin responses, local recycling of ligand-bound receptors in axonal growth cones might also contribute to NGF-dependent functional responses, specifically, local events such as growth cone motility, morphological responses, and guidance. Using a cell-surface biotinylation assay, we did not observe any significant effects of the recycling blocker monensin on levels of TrkA internalized in response to NGF in mass cultures of sympathetic neurons, similar to the results obtained in PC12 cells (Saxena et al., 2005). However, this does not argue against NGF-dependent recycling of internalized TrkA receptors, especially if it occurs in distinct subcellular locations such as axonal processes, which would not have been detected in our assay.

We also observed that NGF treatment results in rapid exocytosis of TrkA receptors to sympathetic growth cones, similar to previous observations for BDNF-induced recruitment of TrkB receptors in hippocampal neurons. BDNF rapidly increased surface TrkB levels within seconds, whereas ligand-induced endocytosis of receptors occurred on a timescale of minutes (Haapasalo et al., 2002). More quantitative kinetic analyses combined with live imaging will be required to temporally separate the two events occurring locally at the sympathetic growth cones. However, it seems reasonable to propose that, like BDNF, NGF rapidly recruits TrkA receptors from a local intracellular pool within axon terminals, followed by endocytosis of active receptors. Because exocytosis of TrkB receptors has been demonstrated to be regulated by activity and intracellular second messengers such as Ca$^{2+}$ and cAMP (Meyer-Franke et al., 1998; Du et al., 2000), it will also be important to define whether similar mechanisms underlie the transcytosis and local exocytosis of TrkA receptors. Target-derived NGF initiates a positive feedback loop by inducing expression of TrkA receptors to enhance the magnitude and duration of prosurvival signaling during neuronal competition for survival (Deppmann et al., 2008). Our results suggest an additional positive feedback mechanism in which limiting concentrations of target-derived neurotrophins rapidly recruit their receptors to axons via transcytosis and promote their exocytosis into growth cones, to amplify neuronal responsiveness and gain a competitive advantage. In this trafficking-based feedback model, the first axons to reach NGF-expressing targets would not only gain access to limiting levels of NGF but also propagate a retrograde signal to neuronal cell bodies to anterogradely transport more receptors to nerve terminals via transcytosis. To integrate our observations, we propose a model in which an initial response of axons to NGF triggers a transient increase in surface TrkA levels through local exocytosis from intracellular receptor pools, followed by endocytosis and retrograde transport of activated receptors in nerve terminals. Subsequently, retrograde NGF signaling enhances anterograde delivery of additional TrkA receptors by drawing on a larger intracellular pool residing in cell bodies (Fig. 6). In future studies, it will be important to define the nature of the retrograde signal that regulates anterograde Trk trafficking, whether it is signaling-endosome-based or mediated by long-range calcium signaling from growth cones to neuronal soma as reported recently for Slit-dependent migration of cerebellar granule cells (Guan et al., 2007), and how this signal is propagated to mature Trk receptors residing on neuronal soma surfaces.

In a broader context, anterograde transcytosis and membrane trafficking from recycling endosomes might also be of relevance to axons navigating through gradients of guidance cues. Axons, during pathfinding, undergo a process of adaptation by consecutive cycles of endocytosis-dependent desensitization and local protein synthesis-mediated resensitization (Ming et al., 2002; Piper et al., 2005). This adaptational mechanism is believed to reset the sensitivity of growth cones to guidance cues as they move up a gradient of an extracellular ligand. Our results show that blockade of Trk receptor recycling attenuates neuronal sensitivity to NGF. Thus, membrane trafficking from recycling endosomes might work in parallel with local protein synthesis, to rapidly mobilize trophic and guidance receptors to axon growth cones, thus allowing them to dynamically interpret gradients of developmental cues.

**References**


