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Lysosomal Activity Associated with Developmental Axon Pruning

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Clearance of cellular debris is a critical feature of the developing nervous system, as evidenced by the severe neurological consequences of lysosomal storage diseases in children. An important developmental process, which generates considerable cellular debris, is synapse elimination, in which many axonal branches are pruned. The fate of these pruned branches is not known. Here, we investigate the role of lysosomal activity in neurons and glia in the removal of axon branches during early postnatal life. Using a probe for lysosomal activity, we observed robust staining associated with retreating motor axons. Lysosomal function was involved in axon removal because retreating axons were cleared more slowly in a mouse model of a lysosomal storage disease. In addition, we found lysosomal activity in the cerebellum at the time of, and at sites where, climbing fibers are eliminated. We propose that lysosomal activity is a central feature of synapse elimination. Moreover, staining for lysosomal activity may serve as a marker for regions of the developing nervous system undergoing axon pruning.

Key words: synapse elimination; lysosome; autophagy; degradation; axon pruning; retraction bulb

Introduction
Clearance of cellular debris is an important aspect of cellular homeostasis. Through normal cellular metabolism, there is a continual buildup of byproducts that are removed via degradative pathways. Most cells in multicellular organisms undergo continual renewal such that old cells die and are replaced by new ones, meaning that the amount of material accumulated in a cell is typically small. Most neurons, however, are postmitotic and do not divide for years or even decades. They are thus particularly vulnerable to the deleterious effects of debris accumulation, and many lysosomal storage disorders manifest with neurological symptoms, suggesting that nervous tissue is especially susceptible to buildup of storage material (Nixon and Cataldo, 1993). Indeed a number of studies suggest that disruption of axonal degradative pathways is linked to axonopathies and neurodegeneration (Yue et al., 2008). Problems with glial digestion pathways, although less frequently described, have been linked to demyelination (Fortun et al., 2003).

In the developing nervous system, synapse elimination involves a massive loss and eventual disappearance of cellular material. This developmental reorganization causes a large majority of nascent synaptic terminals and their associated axonal branches to be removed. In the peripheral nervous system, synapse elimination has been documented in early postnatal life at the neuromuscular junction (Redfern, 1970) and in autonomic ganglia (Lichtman, 1977; Purves and Lichtman, 1980). At the same developmental stages, analogous reorganizations have been found to occur in the cerebellum (Crepel et al., 1976; Lohof et al., 1996), the retinogeniculate system (Chen and Regher, 2000), and the somatosensory system (Arsenault and Zhang, 2006), suggesting that there may be a common mechanism of axon pruning throughout the peripheral nervous system and CNS. Because of the accessibility and large size of neuromuscular junctions, synapse elimination has been studied most extensively in muscle. Time-lapse imaging and serial electron microscopy at the neuromuscular junction shows that retracting motor axon branches are pruned by a shedding process in which axonal fragments (“axosomes”) trail along the path of retreat (Bishop et al., 2004). Electron microscopy shows that these axosomes are engulfed by Schwann cells sheathing the retracting motor axon, but the fate of these axonal remnants within the glial cells remains unclear. The large debris load that glia have to accommodate in early postnatal life suggests that this developmental stage would be particularly sensitive to the effects of disorders in debris clearance.

Here we examine the role of the lysosomal pathway in glia and neurons during the developmental phase, when naturally occur-
ring synapse elimination takes place. We show that regions of the nervous system undergoing synapse elimination can be highlighted with a vital marker that labels late endosomes and lysosomes. We also demonstrate that in a molecular disorder of debris clearance (Cao et al., 2006), retracting axons are removed more slowly. We further provide evidence that by highlighting the lysosomal pathway in the developing brain, regions of axon pruning in the CNS can be identified.

Materials and Methods

Transgenic mice. We used Thy-1 YFP-16 transgenic mice to visualize motor axons as described previously (Feng et al., 2000). Double-transgenic mice were generated for some experiments by crossing Thy-1 CFP-5 mice with S100-GFP transgenic mice to obtain spectrally distinct axonal and Schwann cell labeling (Zuo et al., 2004). GFP-LC3 transgenic (Mizushima et al., 2004) and Cln3Δex7/8 knock-in mouse tissues (Cotman et al., 2002) were received from S.L.C. The Cln3Δex7/8 mice have the most common human Cln3 mutation knocked into the mouse genome.

For studying climbing fibers in the cerebellum, we crossed Thy-1 CFP-5 mice with Thy-1 YFP-16 mice to visualize Purkinje cells and climbing fibers, respectively. Of note, a very small subset of climbing fibers express YFP in the Thy-1 YFP-16 line; thus climbing fiber branches innervating adjacent Purkinje cells are likely to originate from the same climbing fiber axon. Thy-1 CFP-5 X S100-YFP double-transgenic mice were crossed to obtain parallel labeling of Purkinje cells and Bergmann glia. Brainbow line I (Livet et al., 2007) crossed with Cre-ER mice were used to obtain membrane-labeled Bergmann glial expression. Tamoxifen (50 μg) was injected intraperitoneally in postnatal day 4 (P4) pups.

Ex vivo imaging in nerve–muscle explants. We used tissue from Thy-1 YFP-16 or double-transgenic mice (P8–P18) as described previously (Bishop et al., 2004). The triangularis sterni muscle and its innervating nerves were dissected in oxygenated (95% O2/5% CO2) Neurobasal A medium (Invitrogen) at room temperature. The tissue was then pinned into a Sylgard-coated 3.5 cm dish. LysoTracker Red DND-99 (1:5000; Invitrogen) in oxygenated Neurobasal A medium was added to the explant dish and incubated for 2–3 min while being continuously perfused with oxygenated medium. The LysoTracker Red-D99-containing medium was then washed out with new oxygenated Neurobasal A medium to remove excess dye. The tissue was kept heated to 33–35°C and continuously perfused with oxygenated Neurobasal A medium at a perfusion rate of 0.5–1.0 ml/min during imaging. Ex vivo imaging was continued for up to 5–6 h, while the tissue remained healthy and no signs of Wallerian degeneration were noted [for details see Bishop et al. (2004)].

Time-lapse images were taken with a wide-field Olympus microscope equipped with a 20×/0.5 numerical aperture (NA) water and a 100×/1.0 NA water objective lens, and Retiga EXi CCD camera (Qimaging) controlled by MetaMorph software (Molecular Devices). Time-series images were aligned using Autoquant software (Media Cybernetics) to correct for drifts in the image series. Other time series were acquired on an Olympus FV1000 confocal microscope using the above-mentioned water objectives. Images were edited in Adobe Photoshop. To obtain images free of superimposition, in-focus information was manually extracted from individual images of an image stack. When not illustrating quantitatively accurate results but merely the presence of labeling, we used gamma correction to compress the dynamic range of intensities (for example, see Fig. 1 A).

Confocal imaging and immunostaining. Confocal images of fixed tissue were taken using the Olympus FV1000 microscope equipped with a 20×/0.8 NA, a 40×/1.3 NA, and a 60×/1.45 NA oil objective. Confocal image stacks were acquired sequentially, and images were processed with MetaMorph and Adobe Photoshop.

To obtain colabeling of LysoTracker with anti-LAMP-2 antibody, tissue stained with LysoTracker was immediately fixed in 4% paraformaldehyde, which preserves most large but not all small accumulations of LysoTracker staining. This tissue was subsequently stained with a polyclonal anti-LAMP-2 (1:50–100; Zymed) and a Cy5-labeled goat anti-rabbit secondary antibody (1:1000; Invitrogen).

For visualizing axons and synapses in Cln3Δex7/8 knock-in mice (P5, P10, and P14), the sternomastoid muscle was removed from terminally anesthetized animals and then fixed for 30 min with 4% paraformaldehyde. The tissue was immunostained with mouse monoclonal anti-neurofilament SMI-312 antibody (1:1000; Covance) and mouse monoclonal anti-synapsin antibody (1:1000; Sysmatics) and incubated with Alexa686-labeled goat anti-mouse secondary antibodies to obtain axonal staining. Alexa488-conjugated bungarotoxin (5 μM; Invitrogen) was used to obtain postsynaptic acetylcholine receptor labeling. More than 300 total neuromuscular junctions were counted to calculate the incidence of retracting axons in homozygous Cln3Δex7/8 and wild-type littermate pups at each age.

Iontophoretic labeling and EM reconstruction. After ex vivo imaging, tissues were immersion fixed in 2.5% glutaraldehyde and 2% paraformaldehyde in 0.1 m PBS at pH 7.4. Previously imaged bullys with LysoTracker staining were then targeted for serial electron microscopic reconstruction. To create fiducial markers to locate the bullys, sharp electrodes were filled with 1% solutions of DiI (Invitrogen) in 100% methylene chloride and positioned on muscle fibers near the area of interest. Electrical pulses (200 ms, 1–10 V, 1 Hz) were applied to inject a small DiI crystal into the muscle membrane. Photoconversion of the DiI crystals deposited an electron-dense precipitate on muscle fibers flanking the area of interest. Photoconverted tissue was then osmicated in ferrocyanide-reduced osmium for 1 h, dehydrated in ascending ethanol and propylene oxide, and embedded in EMBed812 (Electron Microscope Sciences). Thin sections (1 μm) were cut and examined until the photoconverted DiI crystals could be located. Serial ultrathin sections (~70 nm) were cut through the region of interest. Sections were counterstained with aqueous uranyl acetate and Reynolds’s lead citrate for 30 min each. Sections were examined at 75 kV using a Hitachi H600 transmission electron microscope. Electron micrographs were captured on film and scanned at a resolution of 1000 ppi. Serial sections were aligned and items of interest were segmented using Reconstruct (http://synapsec.nlm.nih.gov/) (Fiala, 2005).

Cerebellar tissue preparation and LysoTracker quantification. Briefly, pups (P5–P30) were anesthetized with a ketamine and xylazine mixture and decapitated, and transverse cerebellar slices (100–200 μm) were obtained using a Vibratome VT1000S (Leica Microsystems) [for details, see Beierlein and Regel (2006)]. Slices were cut in ice-cold oxygenated sucrose-containing Ringer’s solution (in mM: 81.2 NaCl, 23.4 NaHCO3, 69.9 sucrose, 23.4 glucose, 2.4 KCl, 1.4 NaH2PO4, 6.7 MgCl2, and 0.5 CaCl2). Slices were incubated at 32°C for 30 min in sucrose-containing solution, then transferred for 30 min in 32°C artificial CSF (ACSF) solution (in mM: 125 NaCl, 26 NaHCO3, 2.5 KCl, 1.25 NaH2PO4, 25 glucose, 2 CaCl2, and 1 MgCl2), then incubated in ACSF solution at room temperature for 60 min. LysoTracker Red DND-99 (1:5000; Invitrogen) was added to the slices for 10 min during the incubation at room temperature, and the tissue was subsequently fixed in 4% PFA overnight. All solutions during the slicing and incubating steps were oxygenated (95% O2/5% CO2).

After fixation, the tissue was washed in PBS and mounted for confocal imaging and analysis. Purkinje cells in lobes 3 and 9 were imaged and subsequently analyzed for LysoTracker staining. A single optical section for each Purkinje cell was chosen at random, and the total number of LysoTracker-positive structures exclusively surrounding that Purkinje cell soma but not within the Purkinje cell was counted.

Cerebellum image tracings and reconstructions. Reconstruct program (http://synapsec.nlm.nih.gov/) (Fiala, 2005) was used to trace confocal data stacks of fixed LysoTracker-stained cerebellar images. Tracings were performed by hand, and three-dimensional reconstructions were created with Reconstruct.

Results

Acidic organelles associated with neuromuscular synapse elimination

We studied the fate of eliminated axonal branches during development using transgenic mice that express fluorescent proteins in the cytoplasm of axons (Feng et al., 2000). Motor axon branches were imaged during postnatal life at the time the final transitions from multiple to single innervation of neuromuscular
LysoT-positive staining in the neuromuscular junction. A P12 mouse and one retraction bulb (arrow), which is surrounded by strong LysoT staining (red). Arrowheads point to weak axon removal.

To explore this latter possibility, we developed methods to visualize digestive compartments at sites of developmental motor axon degeneration after axonal trauma (Hirata and Kawabuchi, 2002). These results suggest that the degradative events begin within the axon terminal at singly innervated neuromuscular junctions (Fig. 1A, arrowheads). We therefore suspect that extant LysoTracker-positive organelles can fuse with debris within the distal axon, increasing the volume of the digestive compartment.

We found that LysoTracker staining was present in these developing muscles and was distributed in several different patterns around retreating axon branches. At neuromuscular junctions with a bulb-tipped axon, we often observed LysoTracker staining present in the immediate vicinity surrounding the bulb (Fig. 1A; supplemental Movie 1, available at www.jneurosci.org as supplemental material) (38.0%; n = 11 of 29 bulbs). Less often, we observed staining exclusively within the axoplasm of the axonal bulb (Fig. 1B; supplemental Movie 2, available at www.jneurosci.org as supplemental material) (10.3%; n = 3 of 29 bulbs). Finally, at approximately one-half of the neuromuscular junctions, we observed staining both in and around the retreating axon (Fig. 1C) (51.7%; n = 15 of 29 bulbs). These latter two results were surprising, because lysosomes in neurons are described to be commonly located only in the soma and at nodes of Ranvier (Holtzman and Novikoff, 1965). However, we observed accumulations of LysoTracker-positive material within the axon terminal at singly innervated neuromuscular junctions (Fig. 1A, arrowheads). We next turned to time-lapse imaging of retraction bulbs undergoing axosome shedding to see what became of LysoTracker-positive material around retreating axons (1.69; n = 15) was significantly greater than with nonretreating axons (1.60 μm²; SD = 0.001, t test) at the same developmental age. Furthermore, the LysoTracker-positive material surrounding retraction bulbs often appeared as ring-like structures (Fig. 1D). Within some of these rings, we observed fluorescence derived from the axon that we infer to be axosomes (Bishop et al., 2004); in other cases, the rings surrounded a nonfluorescent core (Fig. 1D, arrowhead vs arrow). It is possible that the nonfluorescent cores were axosomes that were slightly further along a degradative process causing the loss of fluorescence. No rings were seen at any singly innervated developing neuromuscular junctions.

We were interested to learn the temporal sequence of lysosomal activity that gave rise to LysoTracker labeling both within and around retreating axons. In earlier developmental stages (P4–P6), there was a higher incidence of intra-axonal labeling, whereas a few days later (P9–P13), lysosomal activity showed a shift to the extraneuromuscular compartment surrounding retreating axons (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). These results suggest that the degradative events begin within the axon and then are completed within the surrounding glial cells.

Degradative organelles associated with axon retraction

We next turned to time-lapse imaging of retraction bulbs undergoing axosome shedding to see what became of LysoTracker-
ringed axosomes. We observed the fluorescence of ringed axosomes to slowly diminish over 1–2 h (Fig. 2A). The loss in intensity was not explained by photo-bleaching, because within the same field, different axosomes lost fluorescence intensity at different rates. LysoTracker–ringed axosomes lost fluorescent protein intensity faster than axosomes not associated with LysoTracker (Fig. 2A, arrowhead and arrow vs asterisk). This difference suggests that LysoTracker staining is associated with the disappearance of axosomes. LysoTracker labels material solely based on its acidic properties (Zhang et al., 1994). Therefore, to confirm that lysosomal organelles are indeed present at sites of LysoTracker labeling, we labeled motor axons in developing muscles using an antibody to LAMP-2, a lysosomal membrane protein specific for late endosomes and lysosomes (Cuervo and Dice, 1996). LysoTracker-stained compartments, including ring-like structures, were outlined by LAMP-2 staining (Fig. 2B). However, LAMP-2 did not mark all LysoTracker-stained structures, suggesting that LysoTracker also labeled a small subset of acidic structures other than lysosomes. This result is consistent with previous studies showing that LysoTracker labels multivesicular and multilamellar bodies (Watts et al., 2004) and autolysosomes (Wang et al., 2006).

Because we saw LysoTracker within the retreating axons, we explored the possibility that autophagy plays a role in axon pruning. We examined developing transgenic mice that contain green fluorescent protein fused to microtubule-associated protein light-chain 3 (LC3), which is known to accumulate on autophagic membranes (Watts et al., 2004) and to colabel LysoTracker-positive autophagic compartments (Wang et al., 2006). Indeed, we observed evidence of autolysosomes both in retreating axons and the surrounding Schwann cells (Fig. 2C,D). The staining pattern found in GFP-LC3 mice is similar to that obtained with LysoTracker. Together, these results corroborate that LysoTracker highlights heteroplastic as well as autophagic processes during synapse elimination at the neuromuscular junction.

To confirm that degradation was occurring in these LysoTracker-positive compartments, we compared light and electron microscopy images of the same axonal bulb showing LysoTracker staining (Fig. 3A, inset). Both a large spherical (arrowhead) and an oblong-shaped structure (arrow) were LysoTracker positive. We relocated these structures in the electron microscope. The retraction bulb was completely sheathed by a Schwann cell (identified by its basal lamina), as were the LysoTracker-positive compartments. We observed a large electron-dense primary lysosome (Fig. 3B) that aligned with the spherical LysoTracker-positive compartment from the fluorescence image (Fig. 3A, inset, arrowhead). In addition, we observed a vesicle-laden structure that coincided with the location of the other LysoTracker-positive structure (Fig. 3C). Consistent with our light microscopic observations of axonal autophagic processes (Fig. 2C), we observed autolysosomes in retraction bulbs at the ultrastructural level (Fig. 3D–I). These results confirm LysoTracker’s ability to detect both axonal and glial degradative processes during axon pruning.

The sheathing of both bulbs and LysoTracker-positive material within Schwann cells suggests that glia are an important component in the process of axonal removal. Indeed, time-lapse imaging in transgenic mice that express fluorescent proteins in Schwann cells showed directly that axon retreat and axosome shedding occur entirely within the confines of Schwann cells (Fig. 4A; supplemental Movie 3, available at www.jneurosci.org as supplemental material). Importantly, the motor axon’s silhouette remained negative during the period of time-lapse imaging, indicating that no fluorescent glial material (e.g., lysosomes) was transferred back to the axon during this process. Moreover, in double-transgenic mice in which both axons and Schwann cells are labeled, we observed colocalization of LysoTracker and axosomes within Schwann cells (Fig. 4B). This LysoTracker-positive material in glia included ring-like structures (Fig. 4C and see above). Interestingly, in some instances axonal bulbs still connected by a thin stalk were nearly completely ringed by LysoTracker within the Schwann cell (Fig. 4D). Retracting axons with this morphology were previously suggested to be at the stage when formation of a large axosome is imminent (Bishop et al., 2004). A phagocytic role of glial cells therefore seems highly likely. In contrast, however, macrophages [as labeled in CX3CR1 knock-in mice (Jung et al., 2000)], which are also potential phagocytic cells and are present in developing muscle, were not observed in association with retracting axons, suggesting that they have no role in axon removal at the developing neuromuscular junction (supplemental Fig. 2, available at www.jneurosci.org as supplemental material).

Retracting axons disappear more slowly in a mouse model of a lysosomal storage disease

Evidence of autolysosomes in the retreating axons and nearby glial cells (Fig. 2C,D) prompted us to investigate whether disrupt-
In developing skeletal muscle of homozygous Cln3<sup>−/−</sup> mice (Gotman et al., 2002), the incidence of axons in the midst of retracting at P5 and P10 was significantly greater in wild-type littermates (at P5, 41.3%, SD = 14.6, 8 pups vs 17.8%, SD = 4.26, in 5 pups; p < 0.01, t test; and at P10, 9.07%, SD = 2.69, in 8 pups vs 4.50%, SD = 1.46, in 7 pups; p < 0.0005, t test) (Fig. 5A–C; supplemental Fig. 3A, available at www.jneurosci.org as supplemental material). Moreover, sometimes two or three retracting axons (n = 6 in 16 muscles) were observed at a single neuromuscular junction (Fig. 5D), a result never obtained in wild-type animals (n = 0 in 10 muscles). The presence of multiple retreating axons per neuromuscular junction is consistent with the idea that there is a delay in the clearance of axon material in these mice. Because ordinarily only one retreating axon from a neuromuscular junction is observed at any time, it is likely that axons that are removed from a neuromuscular junction do so in a sequential way. In the affected pups, however, the removal process is slowed such that multiple withdrawing axons are present at the same time. Neuromuscular junction structure per se showed no obvious phenotype at P5, P10, and P14. Similarly the degree of multiple innervation (i.e., the number of axons that innervate a synapse) was normal (data not shown). By P14, there was no significant difference in the number of retracting axons, and nearly all neuromuscular junctions were singly innervated as in wild-type littermates (supplemental Fig. 3A, B, available at www.jneurosci.org as supplemental material). These results suggest that although there is no defect in synapse formation or elimination per se, disrupting autophagy leads to a transient delay in axonal branch removal.

**Lysosomal activity associated with synapse elimination in the cerebellum**

We next tested whether lysosomal involvement is a feature of developmental synapse elimination in other regions of the nervous system. We examined LysoTracker staining in the Purkinje cell layer of acutely sliced cerebella of neonatal mice at the time climbing fiber inputs to Purkinje cells are lost (Kano et al., 1997). LysoTracker-positive structures were most prevalent during the second postnatal week (Fig. 6A) (2.5 LysoTracker-positive structures per Purkinje cell at P12; n = 42 total Purkinje cells). In contrast, at both P5 and P30, there were fewer LysoTracker-positive structures (0.26 LysoTracker-positive structures per Purkinje cell at P5, n = 53 Purkinje cells; 0.37 per Purkinje cell at P30, n = 71 total Purkinje cells). The increased density of LysoTracker labeling parallels the time course of climbing fiber synapse elimination, which is largely complete by 3–4 weeks of age (Kano et al., 1997). Moreover, most of the LysoTracker staining was localized to the Purkinje cell layer and the nearby deep region of the molecular layer, consistent with the location of climbing fibers that transiently innervate Purkinje cells during development (Scelfo et al., 2003) (Fig. 6B–F). In several cases, we observed robust LysoTracker staining directly apposed to the site of a climbing fiber that innervated a Purkinje cell (Fig. 6C–C′). In contrast to most of the observed climbing fibers that were smooth in appearance, we identified a subpopulation of climbing fibers during the period of climbing fiber re-
moval that had a fragmented appearance (see supplemental Fig. 4, available at www.jneurosci.org as supplemental material). For example, a typical terminal branch of a “fragmented” climbing fiber was on average made up of six disconnected fluorescent pieces (6.0 pieces; SD = 2.05; n = 12 climbing fibers in 4 pups), whereas normally a terminal branch was continuous, i.e., consisted of less than two pieces (1.41 pieces; SD = 0.65; n = 24 climbing fibers in 6 pups; p < 0.02, t test) (i.e., see examples in Fig. 6D,E). In one case, we fully reconstructed two climbing fibers (one fragmented, one not) that innervated adjacent Purkinje cells (Fig. 6D,E). Given the rarity of fluorescent protein expression in climbing fibers in the mouse line we used (see Materials and Methods), these two climbing fiber branches probably originated from the same neuron. The climbing fiber branch that had a fragmented appearance was associated with fourfold more LysoTracker staining (414.4 μm² total area of LysoTracker staining) than the climbing fiber that was not fragmented (110.9 μm²). A significant difference between the amount of LysoTracker associated with normal appearing and fragmented climbing fibers was a general finding (supplemental Fig. 4C, available at www.jneurosci.org as supplemental material). The LysoTracker staining we observed near Purkinje somata and their proximal dendritic trees often appeared in ring-like shapes (supplemental Fig. 5, available at www.jneurosci.org as supplemental material). These rings are reminiscent of the labeling pattern that we observed at developing neuromuscular junctions during synapse elimination (Figs. 1D, 4C). In the cerebellum, these rings appear to be within Bergmann glia, which intimately wrap both Purkinje cells and climbing fibers (supplemental Fig. 5B,C, available at www.jneurosci.org as supplemental material).

Discussion
We show that the digestive mechanisms known as autophagy and heterophagy are involved in the removal of motor axons during postnatal development of the neuromuscular system. The digestion of axonal remnants induces a local increase of lysosomal activity both within retracting axons and their sheathing glial cells. We found that this activity can be highlighted using LysoTracker, a fluorescent vital dye that is trapped by and fluoresces strongly in acidic compartments such as lysosomes. We also assayed the developing cerebellum at the time that climbing fibers are eliminated. Here too we observed a transient increase in LysoTracker staining that closely paralleled climbing fiber loss. Our findings suggest that probes for increased lysosomal activity may generally mark retracting axons during development.

Our work provides several insights into the process of axon removal during development. First, the presence of LysoTracker surrounding the tips of retracting axons is consonant with previous results of a local piecemeal dismantling of axons by nearby glial cells (Bishop et al., 2004). This idea contrasts with the previous suggestions of wholesale branch degeneration (Rosenthal and Taraskevich, 1977) or absorption of axonal material in a strictly reductive process (Riley, 1981). The upregulation of heterophagic (Figs. 3, 4) and perhaps autophagic (Fig. 2C) mechanisms in glial cells near retracting axons suggests that a significant amount of axonal material is isolated and digested by Schwann cells, rather than integrated into the glial cytoplasm. Our previ-
ous work showed that glial cells near retracting fluorescent axons sometimes became fluorescent themselves, suggesting that at least some soluble material from axosomes may end up in the glial cytoplasm (Bishop et al., 2004). Perhaps different axonal components are processed differently in glia.

Second, some axonal material appears to be digested within the axon itself. We observed accumulations of LysoTracker-positive material inside retracting axons (Fig. 1B), and this was consistent with ultrastructural evidence of axonal autolysosomes (Fig. 3D–I). Axonal autophagy may explain the striking atrophy of axons that often precedes synaptic detachment (Bernstein and Lichtman, 1999). These results thus implicate a role for autophagy in developmental axon remodeling. The link between developmental axonal refinements and autophagy is strengthened by our observations of a transient delay in the clearance of axonal material in homozygous Cln3Δex7/8 mice. Studies show that although lysosomal enzyme activity levels in homozygous Cln3Δex7/8 mice are normal, a molecular defect in membrane-trafficking leads to a partially deficient autophagic pathway (Fig. 5) (Cao et al., 2006). This partial deficiency may explain why synapse elimination in homozygous Cln3Δex7/8 mice is resolved by P14. Furthermore, many lines of evidence suggest that compensatory changes occur when particular degradative pathways are disturbed. These include upregulation of lysosomal enzymes (Mitchison et al., 1999) as well as potentially increasing trafficking of degradative organelles through other uninhibited pathways. For example, studies show that when the ubiquitin-proteasome system is disrupted, autophagy compensates by up-regulating digestion of intracellular protein aggregates (Pandey et al., 2007).

The intracellularly sequestered degradative mechanisms used for axon clearance seem distinct from the extracellular proteolytic mechanisms that appear to play important roles during an earlier stage of synapse elimination when axons are thought to detach from the postsynaptic site (Akaaboune et al., 1998; Nelson et al., 2003; VanSaun et al., 2003). Once axons have detached, they become entirely contained within glial cells and would seem to be less vulnerable to enzymes with extracellular modes of action.

Based on the results presented here, we formulate a model of how developing motor axon branches might be removed. This sequential process involves a reiterating cycle that progresses in a distal-to-proximal direction (Fig. 7A). Once an axon is disconnected from the postsynaptic site, autophagy is initiated at its tip. The axon then begins to shed axosomes, which are digested in the adjacent Schwann cell cytoplasm. These events result in the loss of the most distal segment of the axon, leaving behind a Schwann cell that no longer sheaths an axon. This Schwann cell now situated just distal to the axon tip continues to digest shed axosomes (Fig. 7B). The proximal stump of the axon is sheathed by a different glial cell. Our data suggest that the whole process is then repeated on the more proximal segment. What remains unknown is the fate of the Schwann cells after an axon has retreated. In adult animals, we observe only one chain of Schwann cells leading to a neuromuscular junction. Thus the glia associated with the lost axons must be removed. However, their ultimate fate is not presently known.
A number of lines of evidence suggest that synapse elimination in the peripheral neuromuscular system is analogous to events occurring in the developing CNS (Tapia and Lichtman, 2008). We investigated the possibility that climbing fiber branch removal in the developing cerebellum might occur by mechanisms that are similar to those that remove axons in the neuromuscular system. At precisely the developmental stage and site of climbing fiber elimination, we observed an increase in lysosomal activity in Bergmann glia that was highly reminiscent of the lamellar morphologic aspects of Bergmann glial (Golgi epithelial) cell developments in rats. Anat Embryol 177:183–188.


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