Local Control of Neurofilament Accumulation during Radial Growth of Myelinating Axons In Vivo: Selective Role of Site-specific Phosphorylation

Ivelisse Sánchez,*‡ Linda Hassinger,‡ Ram K. Sihag,‡ Don W. Cleveland,i Panaiyur Mohan,‡ and Ralph A. Nixon*‡§

*Department of Psychiatry and §Program in Neuroscience, Harvard Medical School, Boston, Massachusetts 02115; ‡Laboratories for Molecular Neuroscience, McLean Hospital, Belmont, Massachusetts; and Ludwig Cancer Institute at University of California at San Diego, La Jolla, California 92093

Abstract. The accumulation of neurofilaments required for postnatal radial growth of myelinated axons is controlled regionally along axons by oligodendroglia. Developmentally regulated processes previously suspected of modulating neurofilament number, including heavy neurofilament subunit (NFH) expression, attainment of mature neurofilament subunit stoichiometry, and expansion of interneurofilament spacing cannot be primary determinants of regional accumulation as we show each of these factors precede accumulation by days or weeks. Rather, we find that regional neurofilament accumulation is selectively associated with phosphorylation of a subset of Lys-Ser-Pro (KSP) motifs on heavy neurofilament subunits and medium-size neurofilament subunits (NFMs), rising 50-fold selectively in the expanding portions of optic axons. In mice deleted in NFH, substantial preservation of regional neurofilament accumulation was accompanied by increased levels of the same phosphorylated KSP epitope on NFH. Interruption of oligodendroglial signaling to axons in Shiverer mutant mice, which selectively inhibited this site-specific phosphorylation, reduced regional neurofilament accumulation without affecting other neurofilament properties or aspects of NFH phosphorylation. We conclude that phosphorylation of a specific KSP motif triggered by glia is a key aspect of the regulation of neurofilament number in axons during axonal radial growth.

Key words: axon caliber • axon–glia interactions • oligodendroglia • CNS development • protein phosphorylation

Introduction

After reaching their synaptic targets, axons expand up to 3–10-fold to achieve the large diameters required for the rapid conduction of action potentials (Goldberg and Frank, 1979; Graf von Keyserlingk and Schramm, 1984). Expansion is triggered locally when a portion of the axon is wrapped by oligodendrocyte or Schwann cell processes before the elaboration of myelin (Sanchez et al., 1996). Portions of axons or entire axons that remain unmyelinated do not expand (Colello et al., 1994; Nixon et al., 1994). Associated with the process of axon radial growth is a local accumulation of neurofilaments within the expanding region. These 10-nm neuronal intermediate filaments are composed of three subunits designated heavy, medium, and light (NFH,† NFM, and NFL, respectively) according to their respective molecular masses. The requirement for neurofilaments to achieve mature axonal caliber has been clearly demonstrated (Ohara et al., 1993; Eyer and Peterson, 1994), but the mechanisms regulating neurofilament accumulation are unknown.

Neurofilament number within axons is influenced by several factors. After axons make synaptic contacts, the synthesis of neurofilament subunits and other cytoskeletal proteins increases and the rates of slow axonal transport of these proteins decrease severalfold (Hoffman et al., 1983; Willard and Simon, 1983). These changes may raise the levels of many cytoskeletal proteins along the entire length of the axon, but they do not account for the selective expansion of myelinating axonal regions, which

†Abbreviations used in this paper: CNS, central nervous system; ddw, double distilled water; ERK, extracellular signal–regulated kinase; KSP, Lys-Ser-Pro; MBP, myelin basic protein; NFH, heavy neurofilament subunit; NFL, light neurofilament subunit; NFM, medium-size neurofilament subunit; RGC, retinal ganglion cell; TBS, Tris-buffered saline.
achieve diameters and numbers of neurofilaments that may be manyfold larger than those in an immediately adjacent unmyelinated region. Developmentally regulated properties of neurofilaments that might mediate neurofilament accumulation selectively within a discrete region of an axon include the state of phosphorylation and subunit composition as considered briefly below.

During early postnatal development, the COOH-terminal side arm domains of NFH and NFM subunits become extensively phosphorylated within a multiphosphorylation repeat (MPR) region containing the sequence motif Lys-Ser-Pro (KSP) (Carden et al., 1987; Clark and Lee, 1991). In the case of NFH, the >40 KSP repeats present in this region can be separated into two categories, KSPXX and KSPXK (where the final X is any amino acid except lysine), which are believed to be regulated by different protein kinases (Bennett et al., 1994; Elhanany et al., 1994). Several studies suggest that, among numerous candidate neurofilament protein kinases (115 kD, glycogen synthetic kinase [GSK] 3, extracellular signal–regulated kinase [ERK], stress-activated protein kinase, protein kinase K, protein kinase C), CdK-5 preferentially phosphorylates the KSPXK repeats (Hisanaga et al., 1993; Shetty et al., 1993; Guidato et al., 1996; Sun et al., 1996) and has been reported to generate epitopes recognized by the monoclonal antibodies SMI31 and RT97 (Bajaj and Miller, 1997). In contrast, ERKs and GSK appear to regulate KSPXX repeats and generate the SMI34 as well as SMI31 epitopes (Roder and Ingram, 1991; Pant and Veeranna, 1995). Phosphorylation at the COOH-terminal domains of NFH and NFM in vitro straightens individual neurofilaments and promotes their alignment into bundles (Lettrier et al., 1996) and, in vivo, is associated with an increased interneurofilament spacing (Hsieh et al., 1994; Nixon et al., 1994), as NFH and NFM COOH-terminal side arms extend and form crossbridges among neurofilaments and other cytoskeletal elements (Hirokawa et al., 1984; Gotow and Tanaka, 1994; Gotow et al., 1994). Neurofilament transport slows when NFH and NFM are extensively phosphorylated (Watson et al., 1989; Archer et al., 1994). When these subunits are at their highest states of phosphorylation, neurofilaments may stop moving for extremely long periods of time (Lewis and Nixon, 1988), presumably reflecting their integration within a stationary but dynamic cytoskeletal network along axons (for reviews see Hirokawa et al., 1984; Nixon, 1998a,b). Although it is likely that the different properties and behaviors of neurofilaments within axons are independently regulated by the phosphorylation of specific sites within the MPR domain, this possibility has not been previously addressed.

The subunit composition of neurofilaments also influences neurofilament number in axons under some conditions (Cote et al., 1993; Collard et al., 1995; Wong et al., 1995; Xu et al., 1996). The delayed expression of NFH during development occurs during the same period as when slow axonal transport rates decrease and neurofilament levels within axons increase (Hoffman et al., 1983; Willard and Simon, 1983). Moreover, increasing the NFH content of neurofilaments by overexpressing NFH in transgenic mice slows neurofilament transport and raises axonal levels of neurofilaments proximal along axons (Cote et al., 1993; Collard et al., 1995; Marszalek et al., 1996). In both of these situations, the potential contribution of subunit phosphorylation to these effects is unclear. More recently, targeted disruption of neurofilament subunit genes (Elder et al., 1998a; Rao et al., 1998; Zhu et al., 1998) has been used to investigate the roles of each neurofilament subunit in axon radial growth. Ablation of the genes for either NFM or NFL significantly reduced neurofilament numbers and axonal calibers (Ohara et al., 1993; Zhu et al., 1997; Elder et al., 1998a). Neurofilament assembly requires NFL (Ching and Liem, 1993; Lee et al., 1993), and the markedly lowered NFL levels in NFM-deleted mice (Elder et al., 1998b) presumably accounts in part for reduced axon calibers in these mice. Mice lacking NFH have also been generated (Elder et al., 1998b; Rao et al., 1998; Zhu et al., 1998). The small though appreciable effects of this gene deletion on motor axon caliber led some investigators to question the role of NFH in determining axon caliber and, by inference, regulating neurofilament number. However, revealing the function of a protein based solely on the magnitude of phenotypic changes after gene ablation may be problematic because functions of the deleted protein are often compensated by other proteins. In this regard, modest caliber reductions were observed in NFH-deleted mice, but effects of NFH deletion were partially compensated by increases in NFM levels and numbers of microtubules (Rao et al., 1998; Zhu et al., 1998).

To distinguish the roles of subunit stoichiometry and phosphorylation as determinants of regional neurofilament accumulation, we characterized the changes in molecular and structural properties of neurofilaments as they occurred differentially in expanding and unexpanded portions of mouse optic axons during development. We combined analyses of normal retinal ganglion cell development with specific perturbations of neurofilament behavior in NFH-deleted mice and in Shiverer (Shi) mutant mice carrying a mutation of the myelin basic protein (MBP) gene, which interferes with oligodendrocyte maturation (Chernoff, 1981; Shire et al., 1992) and prevents most axons from receiving signals that trigger regional neurofilament accumulation (Sanchez et al., 1996). Our results show that the regional neurofilament accumulation associated with the greatest radial growth of axons begins well after neurofilament subunits achieve mature stoichiometry and, instead, is closely related to the phosphorylation of a subset of KSP sites on the NFH COOH terminus. In the absence of NFH, NFM partially assumes the role of this phosphorylation event during regional neurofilament accumulation.

Materials and Methods

Animals

Breeding of the C57BL/6J mice and tissue dissections were performed as described previously (Nixon and Lewis, 1986; Sanchez et al., 1996) using mice at six postnatal ages (9–120 d). Production of NFH null mice is described (Rao et al., 1998); mice 1.5 yr of age were used.

Electron Microscopy

Mice were anesthetized with halothane gas and the tissue was fixed through intracardial perfusion with 4% paraformaldehyde, 5% glutaraldehyde in 0.1 M PBS, pH 7.4, at room temperature. The optic nerve was dissected and processed as described by Nixon et al. (1994), segmented in 1.2-mm pieces, cleared in propylene oxide, and embedded in Medcast.
(Ted Pella, Inc.). Ultrathin sections were collected and the section containing the initial portion of the retinal excavation was then used as a standard and further ultrathin sections were stained with uranyl acetate and lead citrate and examined in a EX electron microscope (model JEM1200; JEOL) at 80 kV.

**Immunoelectron Microscopy**

Thin sections of epon-embedded tissue were picked up on formvar-coated nickel grids. These sections were etched with 1% periodic acid for 10 min, rinsed in double distilled water (ddw), etched with 3% sodium metaperiodate for 30 min, and rinsed in ddw, followed by 0.05 M Tris-buffered saline (TBS), pH 7.6, for 5 min. Sections were then blocked with 20% normal goat serum in TBS for 60 min at room temperature and then incubated overnight at room temperature in a moist chamber with primary antibody (RT97) diluted at 1:5,000 in TBS containing 1% normal goat serum, 1% BSA, and 0.1% Triton X-100. The sections were washed sequentially in TBS, pH 7.6, for 5 min, in TBS, pH 8.2, for 5 min, and in TBS, pH 8.2, with 0.5% polyethylene glycol for 5 min. Sections were then incubated in 5-nm gold conjugate IgG diluted to 1:25 in TBS containing polyethylene glycol, pH 8.2, for 2 h at room temperature, followed by rinses, first in TBS then in ddw. Finally, the sections were counterstained in uranyl acetate and lead citrate and examined with an EM microscope (model JEM1200; JEOL) with an AMT digital camera.

**Determination of Axonal Diameter Sizes and Quantitation of Neurofilaments**

Axonal diameters were determined from 1,561–4,919 axons at the 50- and 700-μm levels from each of the six postnatal ages analyzed (P9, 12, 16, 21, 30, and 60) and from RT97-deleted and control mice aged 1.5–1.7 yr old. Neurofilament and microtubule numbers in each optic nerve were determined from electron micrographs at the 50- and 700-μm levels by counting 600–1,185 axon profiles representative of the range in calibers in the total axonal population to accurately reflect absolute numbers of neurofilaments in a population of axons heterogeneous in caliber. Values were expressed as number of neurofilaments per 1,000 axons as described previously (Sanchez et al., 1996).

**SDS-PAGE, Immunoblotting, and Two-dimensional Gel Electrophoresis**

The most proximal 2.2-mm segment of the optic nerve from C57Bl/6J mice at postnatal ages of 4 through adult (120 d) was removed at 4°C after mice were killed by cervical dislocation. Segments from eight mice of a given age were pooled and stored at −70°C. The optic nerve cytoskeleton proteins were obtained from pooled segments for each given age. All subunit levels in developing mice were measured based on the standard optic nerve length. In studies of adult NFH-null and Shiverer mice, we used the standard 9-mm length of optic pathway used in previous studies (Nixon and Logvienko, 1988), which extended from the eye to the optic tract. The tissue was homogenized as shown by Chiu and Norton (1982) with 50 mM Tris-HCl, pH 6.8, buffer containing 1% Triton, 50 mM NaF, 5 mM NaVO₃, 10 μM genistein, 0.25 M NaCl, 50 μg/ml leupeptin, 0.1% aprotinin, and 2 mM PMSF. The resulting NFH peptides were lyophilized, solubilized in a 20% acetic acid and 5% formic acid solution, and separated on a preparative C18 reverse-phase HPLC column as described previously (Sihag and Nixon, 1989). The 2–80% acetonitrile gradient was run in 45 min and 1-ml/min fractions were collected. The HPLC fractions from 3 to 30 containing the tryptic or TLCK-α-chymotrypsin NFH peptides were lyophilized, solubilized in 50 mM NH₄HCO₃, and either dot-blotted on a 0.22-μm PVDF membrane or pooled to be further affinity-purified, respectively. Immunoreactivity of the dot-blotted peptides was examined with SMI34, SM31, or RT97.

**Affinity Purification of RT97 and SM31 Immunoreactive Peptides**

Cyanobromide-activated Sepharose 4B (BD Pharmingen) was coupled to protein A-purified RT97 or to SM31 antibodies using the protocol recommended by the manufacturer, and the slurry was poured into 1-ml syringe columns. The tryptic and chymotryptic digests of NFH were prepared as described above. Equal aliquots of the digest were passed thrice through the RT97 or SM31 affinity columns. The unbound peptides were removed by several washes with 10 volumes of TBS. The NFH peptides that bound to the RT97 or SM31 affinity columns were eluted with 0.1 M glycine, pH 2.7, and neutralized with 1 M Tris-HCl, pH 8.0. The column fractions were then analyzed by dot blot on PVDF membranes. The immunoreactivity of the eluted peptides from each column was tested against both RT97 and SM31 by using the ECL development system (Amersham Pharmacia Biotech).

**Results**

**Oligodendroglial-triggered Axon Caliber Expansion Coincides with Regional Accumulation of Neurofilaments**

Axons of retinal ganglion cells are unmyelinated within the retina and for a distance of ~100 μm after they converge at the retinal excavation to form the optic nerve (Fig. 1. A and B). Beyond the lamina cribrosa, which is a specialized meshwork of astrocytes located at 100–150 μm from the retinal excavation, >95% of the axons in adult mice are myelinated, larger in caliber, and contain more
neurofilaments (Fig. 1, A and C) (Nixon et al., 1994). During postnatal development, the cross-sectional areas of optic axons at levels beyond the lamina cribrosa increase an additional 200% under the control of signals from oligodendroglial cells (Sanchez et al., 1996). Our results showed that >80% of the total regional caliber growth occurred between 21 and 30 d postnatally (Fig. 1, D and E). Regional caliber expansion is defined as the percent increase in the axonal cross-sectional area at the 700-μm level (or more distal levels) over the area at a level 50 μm (Fig. 1 E) from the retinal excavation (i.e., Area [700 μm – 50 μm/50 μm] × 100).

Before postnatal day 16, neurofilaments increased modestly in number at axonal levels both proximal and distal to the lamina cribrosa without resulting in a regional differential of neurofilaments along these axons (Fig. 1, F and G). After P21, however, neurofilaments robustly accumulated only at levels distal to the lamina cribrosa (Fig. 1, F and G). This resulted in a 250% higher number of neurofilaments beyond the lamina cribrosa (e.g., 700 μm from the retinal excavation) compared with the number at levels proximal to the landmark (e.g., the 50-μm level). We refer to this phenomenon as regional neurofilament accumulation, which is defined as the percent increase in neurofilament number at the 700-μm level (or more distally) over the number at the 50-μm level (i.e., neurofilament number [700 μm – 50 μm/50 μm] × 100).

Attainment of Mature Neurofilament Subunit Stoichiometry and Interneurofilament Spacing Precede Regional Neurofilament Accumulation and Caliber Expansion

To investigate how subunit stoichiometry contributes to regional neurofilament accumulation, the subunit composition of neurofilaments from optic nerves isolated at stages throughout postnatal development was quantitated by densitometric analysis of Western blots. For comparison, we analyzed isolated neurofilaments from the intraretinal portions of optic axons where neurofilaments undergo minimal reorganization (Sanchez et al., 1996). Although the retina is composed of different cell types, the axons and perikarya of retinal ganglion cells (RGCs) contained nearly all of the NFH immunoreactivity as shown by confocal microscopic analysis of whole mount preparations of retina immunostained with anti-NFH antibodies (data not shown). Analysis of the Triton-soluble NFH pool that has been proposed to incorporate into assembled neurofilaments during development (Shea, 1994) revealed very little if any Triton-soluble NFH in the optic nerve after P10, although a pool of Triton-soluble NFH protein persisted in the retina throughout development (Fig. 2 A). NFH levels increased nearly threefold between P9 and P16 along optic axons, including intraretinal portions (Fig. 2 B). Consistent with these results, the ratio of NFH to NFL rose sharply from P6 and peaked by P12 along axons.
NFH Phosphorylation during Development Occurs Sequentially: Regional Neurofilament Accumulation Coincides Selectively with RT97 Phosphoepitope Expression

To investigate the relationship of NFH COOH-terminal phosphorylation to neurofilament organization, we measured the levels of individual NFH phosphoepitopes on neurofilaments before and during the process of caliber expansion and neurofilament accumulation. To distinguish the three NFH phosphoepitopes identified by the monoclonal antibodies SMI34 and SMI31 (Sternberger and Sternberger, 1983), and RT97 (Coleman and Anderton, 1990) by biochemical means, we separated chymotryptic digests of electrophoretically purified NFH from adult optic axon neurofilaments by HPLC and analyzed individual peptide-containing fractions for SMI34, SMI31, and RT97 immunoreactivity. Some fractions containing little or no SMI31 immunoreactivity were strongly SMI34 immunoreactive. Some fractions containing little or no SMI31 immunoreactivity were strongly SMI34 immunoreactive (Fig. 4 A). Also, the RT97 epitope was not detected in fractions that exhibited the highest SMI31 or SMI34 immunoreactivity. To confirm these results, we immunoadsorbed equal aliquots of a tryptic digest of NFH onto either RT97 or SMI31 affinity columns (Fig. 4 B). Slot blot analyses of eluted fractions from the RT97 affinity column revealed that some fractions containing RT97 immunoreactive peptides were not detected by SMI31. By comparison, some of the fractions eluted from the SMI31 affinity column were immunoreactive to SMI31 but not to RT97, indicating that SMI31 does not recognize the RT97 epitope and that the two antibodies recognize phosphoepitopes located on distinct peptide sequences. In addition, immunoblots of the total NFH subunit pool separated by two-dimensional electrophoresis were probed with SMI33, a monoclonal antibody recognizing NFH independently of its phosphorylation state. These blots show that the adult optic nerve contains multiple isoforms of NFH, spanning a pH range of 7.1 to 5.1 (Fig. 4 C). SMI31 and SMI34 recognized NFH isoforms over a wide pH range (pH 6.8 to 5.1; not shown); however, RT97 cross-reacted only with the most acidic of these isoforms (pH 5.9 to 5.1), in agreement with previous findings suggesting that RT97 recognizes an epitope that is present only in highly phosphorylated NFH peptides (Coleman and Anderton, 1990).
neurofilament spacing was observed only at 700 μm (P9 and P16; thereafter, spacing remained unchanged (Fig. 5 D)). By contrast, the RT97 phosphoepitope remained at negligible levels in the retina and was first detected in optic axons only after the third postnatal week (Fig. 5 B), increasing linearly in amount until P120. Its late appearance was not due to lower expression levels of NFH during early development because equal amounts of NFH protein from P30 and P120 optic nerves yielded the expected 2.5-fold increase in RT97 immunoreactivity (Fig. 5 C). The appearance and rise of RT97 immunoreactivity between P21 and P120 coincided with the onset and rise of regional neurofilament accumulation (compare Figs. 1 G and 5 B).

Preservation of Regional Neurofilament Accumulation in NFH-deleted Mice Is Accompanied by Selectively Increased Generation of the RT97 Phosphoepitope on NFM

Because the small phenotypic effects of NFH gene ablation in peripheral motor neurons (Elder et al., 1998b; Rao et al., 1998; Zhu et al., 1998) have raised questions about the importance of NFH to neurofilament number specification, we investigated whether or not NFM assumes important aspects of NFH function in NFH-deleted mice. The cross-sectional areas of optic axons in NFH-null mice were reduced 12% (0.302 ± 0.013 vs. 0.265 ± 0.020 μm², P < 0.01) or 16% (0.560 ± 0.035 vs. 0.438 ± 0.024 μm², P < 0.01) at the 50- and 700-μm levels, respectively, reflecting mainly a reduced proportion of the largest caliber axons (Fig. 6 A), as previously seen for motor axons (Elder et al., 1998b). NFH gene deletion had relatively small effects on regional axonal expansion (Fig. 6 B) or regional accumulation of neurofilaments as defined in this study (P = 0.05) (Fig. 6 B). NFM levels were not elevated in optic nerves of NFH-deleted mice (Fig. 6 C) in contrast to motor neurons (data not shown) (Rao et al., 1998). Similarly, the levels of SMI31 immunoreactivity on this subunit were unchanged and no generation of the SMI34 epitope was evident on NFM (Fig. 6 C). By contrast, RT97 immunoreactivity on NFM was substantially increased in NFH-deleted mice (Fig. 6 D). Because RT97 detects a series of NFH breakdown products that extend below the molecular mass range of NFM (Fig. 6 D), one-dimensional gels were inadequate to discriminate accurately the contributions of RT97 on NFH and NFM in the optic axons of wild-type mice. This increase was more clearly seen in peripheral nerves where NFH breakdown products were less prominent (inset, Fig. 6 E). Two-dimensional polyacrylamide gel analyses (Fig. 6 E) of optic nerve did partially separate NFM from NFH breakdown products and showed that RT97 immunoreactivity associated with NFM was elevated in NFH-null mice. Little apparent RT97 labeling of NFM was evident in NFH−/− mice when NFH breakdown products were largely separated by isoelectric focusing from the region of the gel containing NFM. The position of NFM relative to NFH was confirmed by reprobing the blot in Fig. 6, D and E,
with RM04. The RT97 immunoreactive signal associated with NFM was estimated to be 5–10% of the signal associated with NFH, based on multiple two-dimensional immunoblots of optic axons developed for equivalent times or on one-dimensional blots of peripheral nerve. Other evidence implies that a relatively small proportion of the total RT97 immunoreactivity on NFH or on NFM in NFH-null mice may be sufficient to establish normal regional neurofilament accumulation. Neurofilament accumulation at 700 μm had plateaued by P30 (see Fig. 1F), even though RT97 levels were only one third of the level at P120 (see Fig. 5B). Moreover, RT97 levels increase even further with age (data not shown), indicating that the level of RT97 associated with the phase of rapid neurofilament accumulation may be a very small fraction of the RT97 level on NFH in the mature adult. A similar effect on NFM phosphorylation was seen in the spinal cord (data not shown) where we also confirmed a previously observed modest increase (less than twofold) in NFM levels (Rao et al., 1998). Thus, when NFH is deleted, at least one of its characteristic properties, the RT97 phosphoepitope, is partially assumed by NFM.

**Figure 4.** SMI31 and RT97 are distinct NFH phosphoepitopes. (A) NFH peptides after chymotryptic digestion were separated by HPLC, and aliquots of all fractions were immunostained in slot blots with SMI34, SMI31, or RT97. Fractions containing high SMI34 or SMI31 signal and no RT97 signal are indicated by arrowheads. (B) NFH tryptic digest peptides after tryptic digestion were affinity purified with either RT97 or SMI31 antibodies. The slot blots of eluted fractions from SMI31 and RT97 affinity columns probed with RT97 antibody (top panel) or with SMI31 antibody (bottom panel). (C) Two-dimensional PAGE immunoblots of NFH from adult optic nerve cytoskeletal fractions probed with SMI33 to detect total NFH or the RT97 phosphoepitope. All blots were also immunostained with an antibody against NFL used as an internal standard to confirm the pH range. RT97 recognizes only the most acidic (highly phosphorylated) NFH isoforms (pH 5.9 to 5.1). IEF, Isoelectric focusing.

**Figure 5.** Sequential appearance of distinct phosphoepitopes on NFH during development. (A) Relative content of different phosphoepitopes associated with NFH in Triton-insoluble cytoskeletal fractions from intraretinal axons (A) and optic nerves (B) from mice at varying postnatal ages. Each data point represents the immunoreactivity level of a given phosphoepitope expressed as the ratio to total NFH immunoreactivity determined with SMI33 antibody. No RT97 immunoreactivity was detected in retinal axons. Each point is the mean and standard error of separate determinations from four different sets of pooled retinas or optic nerve segments (see Materials and Methods). (C) Immunoblots of neurofilament fractions from optic nerves from P16, P30, and P120 mice containing equal amounts of NFH protein probed with SMI33 (left) and a duplicate blot probed with RT97 antibodies (right). (D) The establishment of regional differences along axons in the content of individual NFH phosphoepitopes during postnatal development. To emphasize regional differences, the content of SMI31 or SMI34 in neurofilaments is expressed as a ratio of the optic nerve (optic axons) to retina (intraretinal RGC axons) values. Values >1 reflect a relative increase in content in the optic nerves. RT97 is not included in D because it was not detectable in intraretinal axons.
Selectively Reduced RT97 Phosphoepitope Levels Are Associated with Decreased Regional Neurofilament Accumulation in Myelin-deficient Shiverer Mutant Mice

Mutations in the MBP gene in Shiverer mice (shi/shi) inhibit oligodendroglial investment of most axons and prevent them from receiving signals that trigger regional accumulation of neurofilaments (Sanchez et al., 1996), although neurofilaments in these axons achieve normal intermediate filament spacing (Fig. 7 A). The percentage of axons that do become invested by oligodendroglial cells equals the extent of regional neurofilament accumulation observed (~35% of normal accumulation) (Sanchez et al., 1996) (Fig. 7 B). Thus, the Shiverer mice provided a suitable model to establish whether or not RT97 phosphoepitope expression may be selectively involved in the glial signaling process that regulates regional neurofilament accumulation. Relative levels of the SMI34, SMI31, and RT97 phosphoepitopes were quantitated in NFH subunits of axonal neurofilament fractions from optic nerves of Shiverer mice lacking the MBP gene (shi/shi) (Shine et al., 1992) and mice expressing the normal level of MBP protein (Fig. 7 C). Neurofilaments from shi/shi optic axons contained 32% of the RT97 immunoreactivity detected in heterozygous or wild-type mice. Two-dimensional immunoblots of optic nerves from NFH+/+ (D) and NFH−/− (E) mice stained with RT97 demonstrated the absence of NFH and presence of RT97-positive NFM in NFH−/− mice. The identity and precise position of NFM in D (indicated by oval) were confirmed by reprobing the blots with RM04 (data not shown). The smaller quantity of NFH breakdown products in sciatic nerve samples allowed increases of RT97 on NFM in null mice to be appreciated (E, inset).
whereas the ~65% of axons lacking myelin or glial investment were completely unlabeled.

**Discussion**

Changes in NFH expression, neurofilament subunit stoichiometry, or interneurofilament spacing have previously been proposed to contribute to the regional accumulation of neurofilaments and expansion of myelinated axons (Lee et al., 1993; Nixon, 1998a); however, we establish here that such changes in neurofilament properties largely precede these growth processes. Like neurofilament number, interneurofilament spacing was also modulated regionally although its relationship to myelination and radial growth is complex. Interneurofilament spacing increases along portions of optic axons invested with oligodendrocytes, but a similar change also occurs along distal portions of unmyelinated axons, which do not expand (Nixon et al., 1994; Sanchez et al., 1996). The absence of a clear relationship between interneurofilament spacing and radial growth is also evident in neurofilament transgenic and gene-targeted mice, which exhibit normal filament spacing despite varying changes in caliber (Marszalek et al., 1996; Xu et al., 1996; Elder et al., 1998a,b; Rao et al., 1998; Zhu et al., 1998).

**Role of Subunit Stoichiometry during Regional Neurofilament Accumulation and Caliber Expansion**

We found that as the proportions of NFH in neurofilaments increased during early postnatal development NFM subunit content of neurofilaments decreased commensurately, suggesting that NFH may compete with NFM in binding to NFL or forming 10-nm filaments as proposed previously by Marszalek et al. (1996). These striking changes in neurofilament composition, however, were complete nearly 2 wk before axon calibers expanded regionally. Targeted disruption of the NFH gene diminishes the radial growth of large motor axons to varying degrees (Elder et al., 1998b; Rao et al., 1998; Zhu et al., 1998), although these effects are unexpectedly small given the role suspected for NFH in radial growth. We found comparably small effects on optic axon calibers and showed that regional neurofilament accumulation was also minimally affected. However, the ability of other proteins to assume the functions of a deleted protein is not uncommon in gene ablation studies. In the case of NFH deletion, caliber re-
The subsequent appearance of the SMI31 epitope also preceded regional neurofilament accumulation. Our findings support the hypothesis that SMI31 generation may be related to changes in interneurofilament spacing, in agreement with earlier findings that both interneurofilament spacing and SMI31 immunoreactivity are lower at nodes of Ranvier than along internodal regions (Mata et al., 1992). Both SMI31 immunoreactivity and interneurofilament spacing were normal in Shiverer mutant mice, even though RT97 levels and regional neurofilament accumulation were markedly reduced (see Fig. 7). These observations establish that neither interneurofilament spacing nor phosphorylation represented by the SMI31 epitope regulates regional caliber expansion.

The delayed appearance of RT97 during postnatal development is the only change in neurofilament properties that coincides with the late onset of regional axon caliber expansion during development, parallels the rate of recruitment of axons for radial growth, and segregates selectively within regions of the axon that undergo neurofilament accumulation. Moreover, among the various properties of neurofilaments that change during development, only RT97 levels and regional accumulation are affected when glia-to-axon signaling is blocked in Shiverer mutant mice. When the NFH gene is deleted in mice, the RT97 epitope increases specifically on NFH, allowing this subunit to substitute partially the phosphorylation function of NFH. This result implies that additional tests of the role of KSP phosphorylation on neurofilament function using an in vivo genetic approach will require replacing both NFH and NFM with the appropriate subunits mutagenized at the appropriate sites.

The foregoing results strongly support the conclusion that phosphorylation of a subset of KSP sites is a key aspect of the regulation of neurofilament number in axons during axonal radial growth. RT97 appearance can be viewed as the culmination of a series of developmental events such as NFH expression and proper neurofilament subunit composition and SMI34/SMI31 phosphorylation, which may serve as critical antecedents to the process of neurofilament accumulation and radial growth under normal conditions. Two hypothetical functions of RT97 generation that remain to be tested are its role as an actual trigger of neurofilament accumulation or, alternatively, as a critical neurofilament modification required to stabilize the neurofilament/cytoskeletal after neurofilaments accumulate. Supporting the former possibility are observations that NFH and NFM phosphorylation slows neurofilament transport (Lewis and Nixon, 1988; Archer et al., 1994). Moreover, neurofilaments containing the most highly phosphorylated isoforms of NFH, shown in this study to have the RT97 phosphoepitope, have the slowest net movement (Yabe et al., 2000) and are retained along axons for long periods (Lewis and Nixon, 1988), presumably associated with a nonuniform stationary, but dynamically turning over axonal cytoskeletal network (Hirokawa et al., 1984; Nixon and Logvinenko, 1986; Nixon, 1998a). As proposed previously (Sanchez et al., 1996), abrupt shifts in the number of neurofilaments within a given region of the axon could be induced by local changes in a particular protein kinase or phosphatase activity that modulates the strength of interactions between neurofilaments and the axonal transport machinery, stationary axonal elements, or both. Phosphorylation may also promote stability of these neurofilaments by reducing their susceptibility to

Delayed Appearance of the RT97 Phosphoepitope on NFH Is Selectively Associated with Regional Neurofilament Accumulation

Two distinct NFH phosphorylation events preceded appearance of the RT97 phosphoepitope and were shown to be regulated independently of both RT97-related phosphorylation and regional neurofilament accumulation. The appearance of the SMI34 phosphoepitope coincided with the first detection of NFH protein and reflected phosphorylation of KSPXX motifs along its COOH terminus (Pant and Veeranna, 1995). We observed that NFH subunits initially appearing in the optic nerve had nearly peak levels of this phosphoepitope (i.e., nearly maximal SMI34/SMI33 ratios), indicating that phosphorylation at this site is essentially complete by the time transported neurofilaments reach the level of the optic nerve. This conclusion is consistent with immunocytochemical evidence that SMI34 and Erk1/2, the kinase(s) believed to generate the SMI34 epitope on NFH (Roder and Ingram, 1991; Pant and Veeranna, 1995; Veeranna et al., 1998), are enriched in ganglion cell perikarya (Sanchez, I., L. Hassinger, T. Whee- lock, G. Hauser, and R.A. Nixon. 1996. Soc. Neurosci. 22: 775.7 [Abstr.]). The appearance of SMI34 on NFH subunits still within ganglion cell perikarya suggests a role for this phosphorylation event early after NFH synthesis.

Protections appear to have been offset by several compensatory mechanisms in different fiber systems. Microtubules were substantially increased (Rao et al., 1998; Zhu et al., 1998) and, in one mouse model where microtubule increases could not be demonstrated, caliber reductions were greater (Elder et al., 1998b). The loss of NFH was also accompanied by elevated NFM levels (Rao et al., 1998; Zhu et al., 1998) and possibly by greater NFM phosphorylation (Zhu et al., 1998), although, in two studies, phosphate addition to known sites on NFM (e.g., SMI31) was not increased (Elder et al., 1998b; Rao et al., 1998). The NFM compensatory response is significant in light of evidence that NFM is required to achieve normal axon caliber (Elder et al., 1998a). What is now clear from our studies of optic axons from NFH-deleted mice is that preservation of the capacity to accumulate neurofilaments regionally is accompanied by the selective appearance of the RT97 phosphoepitope on NFM, which is normally an NFH-specific phosphorylation event. Moreover, the normal 2:1 stoichiometry of NFM to NFH (Nixon and Lewis, 1986) and the increase in NFM in some fiber tracts (Rao et al., 1998; our findings) results in a greater capacity of NFM to rescue NFH function in NFH-deleted mice. Even though levels of RT97 signal on NFM are lower than those on NFH, our evidence indicates that only a small proportion of the total RT97 epitope level is normally present during the period of robust neurofilament accumulation. Levels of RT97 are undetectable in Shiverer axons that are not invested with an oligodendroglial cell, whereas the smaller population of axons with primitive myelin sheaths had normal or near normal levels of RT97. These findings, therefore, suggest that axons are able to compensate partially for NFH's functions and highlight a particularly critical role of NFM and phosphorylation generating the RT97 epitope.
proteases (Goldstein et al., 1987; Pant, 1988). The presence of the RT97 phosphopeptidase on tau (Brion et al., 1993) and microtubule-associated protein 1B (Johnstone et al., 1997) during early development suggests that other potentially interacting cytoskeletal elements regulated by RT97-generating protein kinase(s) may also contribute to regional maturation of the axonal cytoskeleton. Characterizing the intercellular and intraneuronal mechanisms regulating this phosphorylation event should help to reveal how different patterns of pathologic neurofilament accumulation arise in human neurodegenerative diseases.

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