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Coordinate control of axon defasciculation and myelination by laminin-2 and -8

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Schwann cells form basal laminae (BLs) containing laminin-2 (Ln-2; heterotrimer α2β1γ1) and Ln-8 (α4β1γ1). Loss of Ln-2 in humans and mice carrying α2-chain mutations prevents developing Schwann cells from fully defasciculating axons, resulting in partial amyelination. The principal pathogenic mechanism is thought to derive from structural defects in Schwann cell BLs, which Ln-2 scaffolds. However, we found loss of Ln-8 caused partial amyelination in mice without affecting BL structure or Ln-2 levels. Combined Ln-2/Ln-8 deficiency caused nearly complete amyelination, revealing Ln-2 and Ln-8 together have a dominant role in defasciculation, and that Ln-8 promotes myelination without BLs. Transgenic Ln-10 (α5β1γ1) expression also promoted myelination without BL formation. Rather than BL structure, we found Ln-2 and -8 were specifically required for the increased perinatal Schwann cell proliferation that attends myelination. Purified Ln-2 and -8 directly enhanced in vitro Schwann cell proliferation in collaboration with autocrine factors, suggesting Lns control the onset of myelination by modulating responses to mitogens in vivo.

Introduction

Myelin increases the speed of neural conduction in thin axons. Defects in myelination cause debilitating loss of function in a variety of congenital and acquired neurological disorders. Mechanisms coordinating myelination in the peripheral nervous system are poorly understood, despite descriptions of cellular events (Martin and Webster, 1973; Webster et al., 1973) and the identification of molecular cues to developing Schwann cells (Mirskey et al., 2002). We show that two members of the laminin (Ln) family of glycoproteins act in concert to regulate the onset of myelination in peripheral nerves.

Peripheral myelination is a concerted process in which Schwann cell proliferation, axon defasciculation, and myelin assembly overlap (Webster, 1971; Martin and Webster, 1973; Webster et al., 1973; Stewart et al., 1993). Premyelinating Schwann cells cover fascicles of cotargeted axons. Their proliferation rate initially matches axonal growth, but increases during myelination to supply Schwann cells for individual axons, at perinatal ages in rodents. Progeny invade fascicles after longitudinal division, which increases Schwann cell density along subsets of axons. Invading cells often transiently ensheath several axons, but retract all but one process and myelinate a single axon. Recurrence of these events ultimately reduces fascicles to axons lacking promyelinating signals, which are defasciculated but remain unmyelinated by the final Schwann cell progeny.

Webster described the progressive defasciculation and myelination of peripheral axons as radial sorting, and proposed that Schwann cell proliferation is intimately involved in the commitment of longitudinal cohorts to defasciculate and ensheath subjacent axons (Webster, 1971; Martin and Webster, 1973; Webster et al., 1973). Although neuregulins have been identified as key signals for Schwann cell proliferation (Garratt et al., 2000), molecular mechanisms that accelerate perinatal proliferation and propel radial sorting are not known.

The one factor known to have specific roles in radial sorting is Ln-2 (merosin), a major component of the Schwann cell surface basal lamina (BL). Lns comprise a family of αβγ heterotrimers. Loss of Ln-2 through mutations in the α2 chain causes a complex neuromuscular disease including peripheral dysmyelination. In the most studied dy and dy2J strains of Ln-2 mutant mice, peripheral nerves contain bundles of unsheathed axons that resemble embryonic fascicles (Bradley and Jenkinson, 1973; Bisceo et al., 1974). This unique pattern of dysmyelination...
presumably represents incomplete radial sorting and has therefore been termed “amyelination.”

Mechanistic hypotheses for amyelination presume endoneurial BLs are necessary for Schwann cell motility and/or differentiation during rapid remodeling (Madrid et al., 1975; Bunge, 1993; Feltri et al., 2002; Chen and Strickland, 2003). Lns that self-polymerize, including Ln-2, are the key structural component of BLs (Yurchenco et al., 2004), and Ln-2–deficient Schwann cells form patchy, discontinuous BLs (Madrid et al., 1975). However, only spinal roots and cranial nerves are severely amyelinated in dy and dy2J mice; sciatic nerves are partially affected and brachial nerves are nearly normal (Bradley and Jenkinson, 1975; Stirling, 1975; Weinberg et al., 1975). One possibility is that BL structure and Ln have limited roles in radial sorting, only critical in large nerves. Alternatively, loss of Ln-2 may be partially compensated by isoforms containing the α1, α4, and α5 chains. Ln α1 is absent in normal nerves, but is expressed in dy2J sciatic nerves; lack of α1 expression in dy2J spinal roots may account for severe amyelination there (Previtali et al., 2003b). Ln α5 is selectively expressed in roots (Nakagawa et al., 2001), which could interfere with α1-Ln heterotrimer assembly in dy2J. Ln α4 is normally low in mature nerves, but is up-regulated in developing nerves and α2-deficient nerves (Patton et al., 1997, 1999; Nakagawa et al., 2001). Targeted deletion of the Ln α1 chain causes more widespread peripheral dysmyelination than occurs in dy mice, consistent with roles for multiple isoforms (Chen and Strickland, 2003). Here, we address independent and combined roles of Lns containing the α2, α4, and α5 chains.

Results

Neuromuscular dysfunction and peripheral neuropathy

When lifted by the tail, Ln α4-deficient mice (Lama4<sup>−/−</sup>) retracted hindlimbs toward the body, with toes clenched (Fig. 1 b). In contrast, normal and heterozygous Lama4<sup>+/−</sup> littermates extended limbs downward, potentially minimizing fall injuries (Fig. 1 a). Lama4<sup>−/−</sup> hindlimb retractions often progressed to rigid rearward extension (Fig. 1 c), but ceased upon landing. Forelimbs were unaffected. Similar suspension-induced hindlimb retraction was observed in juvenile Lama2<sup>dy2J</sup> (dy2J) mice (Fig. 1 d), before permanent contractures (Fig. 1 e). The overlap in dysfunction suggested Lama4<sup>−/−</sup> might possess an abbreviated form of Ln α2-deficient neuromuscular disease.

Ln α2 and α4 are coexpressed in developing muscles and nerves (Patton et al., 1997). As Lama4<sup>−/−</sup> has no apparent myopathy and limited defects at neuromuscular junctions (Patton et al., 2001), we assessed peripheral myelination (Fig. 1, f–j). Normal nerves are composed of large myelinated axons and thin axons ensheathed by nonmyelinating Schwann cells. In addition to properly myelinated axons, Lama4<sup>−/−</sup> sciatic nerves contained bundles of axons lacking ensheathment. EM confirmed such bundles were largely devoid of Schwann cell processes, and found no solitary naked axons. Premyelinating Schwann cells associated with the large bundles occasionally extended processes between axons or established a promyelinating relationship with a solitary axon (Fig. 1 k; unpublished

Figure 1. Amyelinating peripheral neuropathies in Lama4<sup>−/−</sup> and dy2J mice. (a–d) Overlapping postural defects. When suspended, wild type (a) mice extend limbs downward, whereas Lama4<sup>−/−</sup> mice retract and then extend hindlimbs backward (b and c). dy2J mice retract hindlimbs at juvenile ages (d, 4 wk), before the onset of permanent contractures (e, 3 mo). (f–j) Toluidine blue–stained resin sections of adult control (f), Lama4<sup>−/−</sup> (g and h), and dy2J (i and j) sciatic nerves at low (f, g, and i) and high (h and j) magnification. Bundles of unmyelinated axons are present in mutants, but not controls. (k–o) Electron micrographs show most bundles lack intervening Schwann cell processes. Some Lama4<sup>−/−</sup> Schwann cells along large bundles establish promyelinating relations with single axons (k, asterisk), but usually myelinate small bundles altogether (l and m; Table I). Some polyanxonal myelination included intervening Schwann cell processes (m, arrows), possibly from adjacent cells along the nerve. In dy2J, polyanxonal myelination was rare (o, left), but large rafts of partially defasciculated, unmyelinated, mixed caliber fibers were common (o, right). Bar in f, 38 μm (f, g, and i); 15 μm (h and j); 3 μm (k, l, n, and o); 1.8 μm (m).
data). However, most small bundles were polyaxonally myelinated (Fig. 1, l and m; Table I). Defects in Lama4−/− were remarkably similar to amyelination described in dy and dy2J mice (Bradley and Jenkison, 1973; Biscoe et al., 1974; Weinberg et al., 1975; Okada et al., 1977) (Fig. 1, i and j). Indeed, quantitative analysis of the tibial branch revealed no significant differences between dy2J and Lama4−/− in the number of amyelinated axons or their distribution in bundles (Table I). In both dy2J and Lama4−/−, bundles contained mixed caliber axons. The number of larger axons (minimum diameter ≥ 2 μm) in Lama4−/− bundles (average ± SEM: 292 ± 86/tibial nerve; n = 4) was similar to deficits in myelinated axons. Isolated axons were well myelinated and degenerating myelin figures were absent in both mutants. Finally, although we could not rule out limited central nervous system (CNS) defects in Lama4−/−, we found no CNS amyelination in either mutant (Fig. 1, g and i; insets). Thus, independent genetic lesions in Ln α2 and α4 produce essentially similar amyelinating peripheral neuropathies.

Similarity between Ln α2- and α4-deficient neuropathy included origin and progression (Fig. 2). Most axons in wild-

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**Table I. Amyelination in nerves and roots of Ln α2- and α4-deficient mice**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Wild type</th>
<th>Lama4−/−</th>
<th>Lama2dy2J</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tibial N. (n)</td>
<td>3</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>Myelinated axons</td>
<td>2226 ± 16</td>
<td>1697 ± 49*</td>
<td>1258 ± 16*</td>
</tr>
<tr>
<td>Amyelinated axons</td>
<td>0</td>
<td>2043 ± 458</td>
<td>2380 ± 600</td>
</tr>
<tr>
<td>Bundles</td>
<td>0</td>
<td>31 ± 4</td>
<td>31.0 ± 1.4</td>
</tr>
<tr>
<td>Nonmyelinated bundles</td>
<td>NA</td>
<td>17 ± 3**</td>
<td>28.5 ± 0.7**</td>
</tr>
<tr>
<td>&quot;myelinated&quot; bundles</td>
<td>NA</td>
<td>14 ± 2**</td>
<td>2.5 ± 2.1**</td>
</tr>
<tr>
<td>Axons/bundle (total)</td>
<td>NA</td>
<td>67 ± 16</td>
<td>77 ± 22</td>
</tr>
<tr>
<td>Axons/nonmyelinated bundlea</td>
<td>NA</td>
<td>105 ± 9</td>
<td>80 ± 9</td>
</tr>
<tr>
<td>Axons/myelinated bundlea</td>
<td>NA</td>
<td>18 ± 1</td>
<td>16 ± 5</td>
</tr>
<tr>
<td>V. root (n)</td>
<td>3</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Myelinated axons</td>
<td>760 ± 53</td>
<td>698 ± 56**</td>
<td>32 ± 7**</td>
</tr>
<tr>
<td>Amyelinated axons</td>
<td>0</td>
<td>21 ± 6**</td>
<td>502 ± 42**</td>
</tr>
<tr>
<td>Bundles</td>
<td>0</td>
<td>2 ± 1</td>
<td>10 ± 2</td>
</tr>
<tr>
<td>Axons/bundle (total)</td>
<td>NA</td>
<td>11 ± 1**</td>
<td>49 ± 21**</td>
</tr>
</tbody>
</table>

Values represent mean ± SEM across nerves. *, Different from wild-type value; P < 0.05. **, Difference between mutant values; P < 0.01.

aErrors represent pooled bundles of all nerves.

---

**Figure 2.** Origin and distribution of amyelination in dy2J and Lama4−/− mice. Toluidine blue stained sections from wild-type (a–g), dy2J (h–n), and Lama4−/− (o–u) sciatic nerves at indicated postnatal age, or spinal roots and brachial nerves from 6–9-wk adults. Axon fascicles are sorted by P12 in controls, but persist as amyelinated bundles in mutants. Bundles in year-old mutants are small and surrounded by lightly myelinated fibers (k and l). Spinal roots are severely amyelinated in dy2J (l and m) but not Lama4−/− (s and t). Brachial nerves contain few amyelinated axons in either mutant (n and u). Bar in o, 14 μm (a, h, and o); 8 μm (b, i, and p); 16 μm (c, d, j, k, q, and r); 20 μm (e–g, l–n, and s–u).
type, \( \text{Lama}_2 \) and \( \text{Lama}_4^+/+ \) sciatic nerves were sorted by P5 and myelinated by P12. In \( \text{dy}^2 \)J and \( \text{Lama}_4^+/- \) strains, axon fascicles persisted throughout postnatal development (Fig. 2, h–j and o–q). The proposition that axon bundles in \( \text{dy} \) strains arise through incomplete radial sorting (Bradley and Jenkinson, 1973) is generally accepted, although lack of efficient methods to identify neonatal mutants prevented developmental studies. We confirm that amyelinated bundles are remnants of embryonic fascicles in both \( \text{dy}^2 \)J and \( \text{Lama}_4^+/- \) mice. Amyelination has also been thought permanent (Bradley and Jenkinson, 1973). However, we found only a few, small axon bundles remaining in year-old \( \text{dy}^2 \)J and \( \text{Lama}_4^+/- \) sciatic nerves, each surrounded by lightly myelinated axons (Fig. 2, k and r). As axonopathy was not observed among amyelinated axons, bundles likely erode slowly through continued myelination at their edges (Fig. 1 k). Regardless, \( \alpha_2^+ \) and \( \alpha_4^+/- \) deficient neuropathies did not diverge between onset and old age despite considerable remodeling.

Further comparison revealed two significant differences. First, amyelination is especially severe in \( \text{dy}^2 \)J spinal roots, but was nearly absent from roots in \( \text{Lama}_4^+/- \) (Fig. 2, l, m, s, and t; Table I). Roots are less affected than distal nerves in \( \text{Lama}_4^+/- \). We infer amyelination primarily reflects relative dependence on Ln isoforms, rather than nerve diameter or proximity to spinal origin. Sorting in roots depends strongly on Ln-2 and weakly on Ln-8; distal nerves rely more equally on both. Otherwise, amyelination was distributed similarly in \( \text{Lama}_4^+/- \) and \( \text{dy}^2 \)J; forelimb and intercostal nerves were less affected than hindlimb (Fig. 2, n and u; unpublished data).

Second, polyaxonal myelination (Fig. 1, l, m, and o; Table I) was common in \( \text{Lama}_4^+/- \) but rare in \( \text{dy}^2 \)J, consistent with previous observations in \( \text{dy} \) (Okada et al., 1977). Most
small bundles (≤25 axons) in Lama4−/− tibial nerves were “myelinated.” Conversely, dy2J nerves contained many islands of ensheathed-but-not-myelinated axons, which were rare in Lama4−/− (Fig. 1 o, right). Superficially similar to Remak bundles, islands were less condensed, more numerous, and included mixed caliber axons. They appear to be abnormal transitional structures, intermediate between amylinated bundles and properly myelinated axons, which preferentially appear in dy2J nerves when fascicles contain few axons. Thus, although Ln-2 and -8 both promote axon sorting, they have decidedly unequal roles in roots, and dissimilar roles in the transition from premyelinating to myelinating Schwann cell phenotype.

Redundancy and compensation in the BL
To ask if Ln-2 and -8 independently incorporate into endoneurial BLs, we stained cryostat sections of normal and mutant sciatic nerves with Ln chain–specific antibodies. In normal nerves, Ln α4 was coconcentrated with the α2, β1, and γ1 chains at ab-axonal Schwann cell surfaces; none were detected at axonal surfaces (Fig. 3, d, e, m, and n; unpublished data). Detection of α4 varied considerably with antibody, but endoneurial BLs stained weakly compared with perineurium, as shown previously (Patton et al., 1997; Nakagawa et al., 2001). Loss of α4 did not affect staining for α2 at any postnatal age (Fig. 3 u; unpublished data). Similarly, α4 staining was not decreased in dy2J nerve; indeed, levels increased relative to controls, confirming previous results (Patton et al., 1997, 1999; Nakagawa et al., 2001) with additional reagents. In addition, staining for entactin, perlecan, agrin, and collagen IV was unaffected in either mutant (Fig. 3, g–i, p–r, and y–a; unpublished data). Thus, amylination-inducing mutations in α2 and α4 do not act by inhibiting expression of their counterpart specifically, or disrupting the molecular composition of endoneurial BLs generally. We found no morphological or histological defects in double-heterozygous Lama2<sup>+/−</sup>;Lama4<sup>−/−</sup> offspring (not depicted). Therefore, Ln-2 and Ln-8 each contribute a distinct activity necessary to complete radial sorting.

Next, we asked if myelination achieved in dy2J and Lama4<sup>−/−</sup> reflects partial compensation between Ln-2 and -8, by generating double mutant dy2J/α4null mice (see Materials and methods). dy2J/α4null mice had normal birthweight, fed actively, and responded to stimuli, but were dyskinetic by P14. Adults had splayed stance and strong tremor, moved haltingly, retracted all limbs when suspended, were 50% smaller than normal littermates, and rarely survived 3 mo (Fig. 4, a–f). Peripheral nerves were translucent, rather than the opaque white of myelinated regions of the nervous system. Histological sections showed nearly all forelimb and hindlimb axons were concentrated in large amylinated bundles (Fig. 4, g and h), dramatically exceeding defects in the parent dy2J and Lama4<sup>−/−</sup> strains. Premyelinating Schwann cells surrounded bundles, and a few isolated axons were well myelinated, demonstrating neural crest migration and proliferation of Schwann cell precursors was not prevented. CNS myelination appeared unimpaired (Fig. 4 h, inset), showing potential metabolic disorders in these mice do not, per se, prevent formation of myelin. Thus, Ln-2 and -8 act in concert to play a dominant role in the myelination of distal nerve, and each only partially compensates the other’s deficiency.

Remarkably, however, spinal roots in dy2J/α4null mice were completely sorted. Compared with dy2J, the additional...
loss of Ln α4 strongly enhanced radial sorting in dy2J/a4null roots (Fig. 4, i and j; compare with Fig. 2, l and m). Paradoxically, Ln-8 promotes radial sorting in distal dy2J nerves but inhibits sorting in dy2J roots. To explain disparate results in root and distal nerve, we sought differences in Ln receptors and components of the endoneurial matrix that could modulate Schwann cell responses to Ln-2 or -8. We found no reliable differences in receptor components integrin α3, α6, β1, and β1D subunits, α-dystroglycan, β-dystroglycan, or β-sarcoglycan, between normal, dy2J, or Lama4−/− mice (unpublished data), extending results in dy2J and dy3K (Nakagawa et al., 2001; Previtali et al., 2003a,b). Two matrix differences between sciatic nerves and roots are reported; both are Ln α chains. Ln α1, which is absent in normal endoneurium, is expressed in dy2J sciatic nerves, but not their roots (Previtali et al., 2003b). In contrast, Ln α5 is expressed in normal and α2-deficient roots, but not distal nerve (Nakagawa et al., 2001).

Several aspects of Ln α1 expression were inconsistent with roles in radial sorting. First, α1 expression and myelination were poorly correlated in adult dy2J (6–9 wk). α1 was undetectable on many myelinated fibers (Fig. 3 k), and was absent from any brachial or sciatic fibers in 4 of 8 dy2J mice. Several α1 antibodies, which strongly stained CNS pial surfaces included as controls, gave similar results. Second, we were unable to detect α1 in dy2J nerves before P14, when radial sorting has largely ended even in mutants (Fig. 5, d and e; unpublished data). Third, α1 was absent from partially myelinated nerves in Lama4−/− (Fig. 3 t) and α2-null dy3K mice (Fig. 5 f). Fourth, levels of α1 were highest in severely amylmented dy2J/a4null nerves (Fig. 5, g–i), indicating endogenous α1 expression is insufficient for radial sorting. As Ln-1 (α1β1γ1) is elsewhere implicated in BL assembly (Yurchenco et al., 2004), the heterogeneous expression of Ln α1 we observed in dy2J may account for the variable integrity of endoneurial BLs present (but not often acknowledged) in this strain. In dy2J/a4null nerves, α1 was predominantly associated with myelinated fibers (Fig. 5, h and i), many of which contained well-formed BLs (Fig. 4 k). Pre- and promyelinating dy2J/a4null Schwann cells lacked BLs (Fig. 4 k). Thus, Ln α1 may not promote radial sorting because α2-deficient Schwann cells express it after myelination.

Next, we asked if Ln α5 fosters radial sorting in dy2J/a4null nerve roots. Lama5−/− mice die as embryos (Miner et al., 1998), before radial sorting. Therefore, we used a broadly expressed α5 transgene (TgA5) (Kikkawa et al., 2002) to ask if ectopic α5 expression would promote sorting and myelination in the distal portions of dy2J/a4null nerves. Although TgA5 did not increase levels of Ln α5 in wild-type or dy2J endoneurial BLs (unpublished data), possibly due to competition from endogenous α2 and α4 chains for heterotrimer assembly, α5 was readily detected in dy2J/a4null/TgA5 nerves (Fig. 5 k). α5 colocalized with the β1 and γ1 chains, and β2 was ab-
sent (not depicted), indicating ectopic expression of Ln-10 (α5β1γ1). Ln-10 expression was accompanied by suppression of tremor and dyskinesia, and a marked increase in sorting and myelination in distal nerves (Fig. 5, k and l; compare with Fig. 4 g). The results implicate Ln-10 in the sorting and myelination of axons in dy2J/a4null roots, but do not establish whether Ln-10 acts autonomously or depends on the truncated isoform of Ln-2 produced from the dy2J allele of Lama2.

Cell and molecular mechanisms

Amyelination has been thought to derive in large measure from disruption of Schwann cell BLs (Madrid et al., 1975; Bunge et al., 1986; Eldridge et al., 1989; Feltri et al., 2002). However, amyelination in Lama4−/− was not accompanied by disruption of endoneurial BLs, on either myelinating or premyelinating Schwann cells (Fig. 3 d; unpublished data). Moreover, Schwann cells in dy2J/a4null roots ensheathed and myelinated all axons despite lacking even a trace of endoneurial BLs (Fig. 4 l). Promyelinating Schwann cells in dy2J/a4null/TgA5 nerves also lacked BLs (Fig. 5 n). Thus, BLs are neither sufficient (in Lama4−/−) or necessary (in dy2J/a4null) for Schwann cells to complete radial sorting.

Therefore, we considered potential signaling roles for Lns. Several lines of research suggest signaling through Ln receptors might promote Schwann cell proliferation during myelination. Radial sorting is closely coupled to Schwann cell mitosis (Bradley and Asbury, 1970; Webster et al., 1973), amyl elinated regions of Lns α2 mutant nerves have fewer Schwann cells than normal (Bray and Aguayo, 1975; Okada et al., 1976), and Ln-1 promotes Schwann cell proliferation in vitro (Porter et al., 1987). First, we asked if Schwann cell deficits correlate specifically with amyl elination, or with loss of Ln-2 or BL structure. We found Schwann cell deficits specifically accrue from amyl elination axons.

Second, we asked if Schwann cell deficits temporally correlate with radial sorting. Sciatic nerves in Lama2/α4-DKO, dy2J/a4null, and normal littersmates had statistically similar numbers of Schwann cells at E17 (Fig. 6 d). Severe deficits in Lama2/α4-DKO nerves accrued almost entirely between E17.5 and P0. Controls, 12 ± 4 nuclei/field at 1000×; Lama4−/−, 14 ± 5). Thus, Schwann cell deficits specifically accrue from amyl elination axons.

Figure 6. Lns increase perinatal Schwann cell proliferation. (a) Adult Lama4−/− and dy2J/a4null nerves have fewer Schwann cells per unit length than age-matched controls. Bars show nuclei per 8 μm transverse section of tibial branch at medial level; means ± SEM, n mice; 4–9 serial sections/nerve. (b and c) Myelinated fibers teased from wild-type (control) and Lama4−/− tibial nerves have similar Schwann cell density; S100 (green); Hoechst (blue); values in text. (d) Ln-deficient nerves have normal numbers of Schwann cell nuclei at E17.5. Bars show nuclei per 10 μm section of entire medial sciatic nerve, as in a. Combined data from dy2J/a4null (n = 2) and Lama2/α4-DKO (n = 3) are not statistically different from normal littersmates. (e–j) Quadruple-stained sections of P3.5 tibial nerve from a normal (e–g) and Lama2/α4-DKO (h–j) littermate pair: Ki67 (pseudocolored green), Hoechst (blue), integrin α6 (red), and TUNEL (FITC shown in grayscale; g and j). Compared with controls, mutant nerves are thinner and have fewer cell soma, proportionally fewer proliferative (Ki67 labeled) cells, and almost no necrotic cells. Quantitation of total soma (k) and proliferative cell label index (l) in 10 μm tibial nerve sections shows perinatal expansion of Schwann cell populations in normal littersmates (open symbols, bars), but not Lama2/α4-DKO (filled symbols, bars). Proliferation in mutants maintains immature Schwann cell density [numbers per 10 μm length] at embryonic levels during postnatal limb growth. Bar in j, 40 μm in b and c; 27 μm in e–j. Asterisk indicates P < 0.05 for ANOVA comparison between age-matched data sets.
(14.7 ± 8.1%) and P3.5 (73 ± 6.9%) (Fig. 6 k). Moreover, deficits occurred through inadequate proliferation rather than cell death (Fig. 6, e–l). Proliferating cells were identified with antibody to Ki67 (Lalor et al., 1987), and necrotic cells by TUNEL assay. At P3.5, 40% of normal (littermate control) endoneurial cells were Ki67-positive, whereas <20% were labeled in Lna2/a4-DKO pups (compare with values in Fig. 6, k and l). Fewer than 1% of nuclei in normal and mutant nerves were TUNEL stained at any perinatal age (Fig. 6, g and j; unpublished data). Thus, Lna2 and -8 are specifically required for the perinatal increase in Schwann cell proliferation that coincides with radial sorting. They appear dispensable for the proliferation of immature Schwann cells covering fascicles, consistent with EM observations.

To ask if Lna2 and -8 promote proliferation directly, we cultured primary Schwann cells on substrates containing purified isoforms (Fig. 7, a–d). Populations plated at moderate densities on Ln-1, -2, and -8 expanded at similar rates, doubling the rate on uncoated surfaces. When Ln concentration or cell density were limiting, proliferation was significantly faster on Ln-8 than on Ln-1 or -2. These data extend previous studies with Ln-1 (Porter et al., 1987) to suggest that Lna2 and -8 promote Schwann cell proliferation in concert with autocrine growth factors. The results are consistent with the early hypothesis that increasing cell density activates Schwann cells to invade fascicles and ensheath axons (Martin and Webster, 1973).

Lastly, we used adhesion assays to ask if Schwann cells interact with Ln-2 and -8 through distinct or similar receptors (Fig. 7, e–o). Adhesion to purified Ln-1, -2, and -8, but not poly-lysine, was blocked by EDTA, consistent with a role for integrin receptors. Antiserum raised against Ln-8 blocked adhesion to Ln-8 but not Lna2, indicating that binding to Ln-8 relies on distinct epitopes. Finally, adhesion to Ln-8 but not Lna2 was inhibited by function-blocking antibody to integrin α6. The simplest interpretation of these data is that Lna2 and -8 regulate Schwann cells through distinct integrin subtypes. Consistent with this notion, Schwann cell–specific disruption of the integrin β1 gene (Feltri et al., 2002) produced the pattern of amyelination (partial in sciatic nerve; absent in roots) characteristic to Lama4-/-, and distinct from that in dy2J. The combined results suggest Ln-8 (and not Lna2) promotes radial sorting through integrin α6β1.

**Discussion**

We establish a dominant role for Lns in peripheral myelination, identify which of the four isoforms (Ln 1, 2, 8, and 10) expressed by Schwann cells are primarily involved, and clarify mechanisms by which they act. Lna2 and Ln-8 act in concert to increase rates of Schwann cell proliferation at the onset of radial sorting, such that combined Lna2/Ln-8 deficiency prevents radial sorting altogether. Lna2 and -8 also regulate the onset of myelin formation by postmitotic Schwann cells, but through distinct effects on axonal ensheathment. At both steps, their combined activities foster solitary relationships between myelinating Schwann cells and axons (Fig. 8).

**Functions for Lns in nerve development**

Mechanisms underlying the transition from immature Schwann cells surrounding axon fascicles to myelinating and nonmyelinating Schwann cells ensheathing individual axons are not known. Martin and Webster (1973) observed that involution of axons along fascicle edges is immediately preceded by Schwann cell mitosis and an increase in cell density along those axons, and proposed that Schwann cell proliferation plays a primary role in the onset of radial sorting. Indeed, developing Schwann cell populations appear to expand in two
A Perinatal Schwann cell proliferation

B Axonal ensheathment

Figure 8. Proposed roles for Ln-2 and Ln-8 in peripheral nerve development. (A) Developing Schwann cells are regulated by autocrine (1) and axon-derived (2) factors. Ln-2 and -8 are concentrated in Schwann cell BLs, on ab-axonal surfaces (3). Immature Schwann cells initially proliferate to cover axon fascicles, without dependence on Ln-2 or -8. Radial sorting commences as combined Ln-2, Ln-8, and mitogen signals increase Schwann cell proliferation rates and stimulate progeny to envelop subaxcent axons (4). (B) Ln-2 and -8 differentially regulate axonal ensheathment. Ln-8 promotes small ensheathing processes and/or delays myelin formation. Ln-2 promotes onset of myelination and/or inhibits formation of ensheathing processes.

Molecular mechanisms

Ln-2 and -8 do not fully compensate each other’s loss in vivo, and have distinct binding and proliferative activities for Schwann cells in vitro, suggesting distinct receptors mediate their actions. Schwann cells express several Ln-binding integrins and dystroglycan (Pevitari et al., 2001). Early steps of myelination depend greatly on β1-integrins (Fernandez-Valle et al., 1994; Feltri et al., 2002), and not dystroglycan (Saito et al., 2003). Amyelination caused by loss of integrin β1 (partial in distal nerves; nearly absent in roots) now appears to largely phenocopy loss of Ln-8 rather than Ln-2. The major β1-integrin in developing Schwann cells is integrin α6β1. As blocking antibody to integrin α6 inhibited Schwann cell binding to Ln-8 and not Ln-2 (Fig. 7), the simplest interpretation at present is that integrin α6β1 primarily mediates the effects of Ln-8 in radial sorting. Ln-2 engages additional receptors, as adding its loss to α4-deficiency (i.e., in dy2J/α4null and Lama4/α4-DKO) produces amylination far exceeding that in β1-integrin–deficient nerves. Testing in vivo roles for integrin α6β4, expressed by perinatal Schwann cells, may require tissue-specific mutations, as mice lacking these subunits die at birth (Dowling et al., 1996; Georges-Labouesse et al., 1996).

Endoneurial BLs

The idea that Schwann cell BLs are necessary for the proper defasciculation and myelination of axons in developing nerves and Aguayo, 1975; Stirling, 1975; Perkins et al., 1980). However, perinatal proliferation is reportedly normal in mice with Schwann cell–specific loss of Ln γ1 or integrin β1, both of which are amylinated (Feltri et al., 2002; Chen and Strickland, 2003). Interestingly, disruption of floxed Ln γ1 alleles occurred unexpectedly in the Schwann cell lineage at E17, through Cre expression off a CaM kinase II (CaMKII) promoter (Chen and Strickland, 2003). As this coincides with the onset of Ln-dependent proliferation we identify, CaMKII-Cre expression could begin in Ln-dependent progeny. Defects in CaMKII–Cre/Lnγ1-deficient nerves would be consistent with additional roles for Ln-2 and -8 in the invasion of fascicles and/or ensheathment of axons, as suggested previously (Chen and Strickland, 2003). Similarly, β1 integrins could preferentially mediate these later roles (Feltri et al., 2002).

Our results suggest Ln-2 and -8 coordinate axonal ensheathment by promyelinating Schwann cells. Lama4−/− Schwann cells prematurely myelinated bundles of axons, before completing their separation. dy2J Schwann cells lingered between ensheathing multiple axons and myelinating single axons. In principle, both activating and inhibitory mechanisms could underlie these results. For example, Ln-8 may promote the formation of multiple ensheathing Schwann cell processes, or simply retard myelin formation; loss of either function could produce polynodal myelination. Similarly, Ln-2 may promote myelin formation, and/or limit the formation of axon-ensheathing processes. Regardless, the results provide an initial insight into the mystery of how each Schwann cell manages to myelinate a single axon.
develops without BLs, and is almost entirely dependent on Ln-8 to promote BL formation, but is consistent with observations in (ster and Billings, 1972).

in which Schwann cells acquire BLs after myelination (Wetsens brings mammalian myelination into line with amphibians, That BL integrity is irrelevant to the initial myelination of ax-

complete radial sorting nor sufficient to prevent amyelination. (Madrid et al., 1975; Weinberg et al., 1975; Nakagawa et al.,

oning and myelination in

terminal stages whenever possible. C57BL/6J and Lama20def mice were from The Jackson Laboratory. Mice null for Lama2 (Lama20/0) and Lama4 were described previously (Miyagoe et al., 1997; Patton et al., 2001); both were backcrossed five generations to C57BL/6J. Mutant alleles of Lama2 and Lama4 on chromosome 10 were linked by mating heterozy-
gous males (Lama20/0;Lama4/0; or Lama20/0;Lama40/0) to wild-type females and screening offspring for possession of both mutant α2 and α4 alleles (which required paternal recombination). Lamains recombi-
ned with Lama20/0 in 3 of 171 offspring, and with Lama20/0 in 2 of 125 offspring, in agreement with reported genetic distance (Miner et al., 1997). Linkage was confirmed by mating with wild types. Mutants were born at Mendelian frequencies in all founder lines. dy2J/α4null and α2/

4nullDKO mutants did not differ significantly between lines, or from F1 mutants from inter-line crosses in which sites of recombination remain hetero-
ygous, and their data were pooled. Ln α2 and α4 were undetectable in Lna2/4/α4DKO tissues, as in single mutants (Miyagoe et al., 1997; Pat-
ton et al., 2001). A previously described full-length mouse Ln α5 cDNA transgene (Kikawara et al., 2002) prevents embryonic lethality and rescues all known defects in Lama5 mice.

Genotypes were identified by PCR off tootip DNA, using the following sense [5] and antisense [AS] primers. Lama40S: 5’-GGAGCACGTCGCT-
CCACTGTC-3’; AS: 5’-CAACAAATGGTGAAGGTGCCC-3’C. Lama40S:
S: 5’-AGCGATCCTTCCCCCACCC-3’; AS: 5’-GTCAAGGCGTCGC-
CTGCAGACT-3’, in PKG promoter. Lama2α2S: 5’-CCAGATGCTCCA-
CTTGCTC-3’, AS: 5’-CTTCCCTTACCTTGGAAG-3’; Lama2α2S: 5’-
CITTC-
CCAGATGCTCCA-
CTTGCTC-3’, AS: 5’-AGCCATC-
TAGCTTTG-3’. touch-down PCR across the point mutation site (annealing from
65°C to 52°C in 10 cycles plus 2 cycles at 50°C) was followed by Nextel
digestion to produce 164- and 109- bp fragments from dy2J but not wild-
type product. Digestion ambiguities were resolved by HypChat 3, which cuts wild-type but not dy2J product.

Antibodies
Rat mAbs 198 and 200 to Ln α1 were from Lydia Sorokin (Sorokin et al., 1992; Lund University, Lund, Sweden). Rabbit antibodies to Ln α1 and α2 (Rambukkana et al., 1997) were from Peter Yurchenco (Robert Wood Johnson Medical School, Piscataway, NJ); α1-null was generated against EHS Ln-1, affinity purified against E3 fragment, and cross-adsorbed against E8 fragment. mAbs to α2 (4H8-2; Biogene), b1 (MAB1928; CHEMICON International), and γ1 (MAB1914; CHEMICON Interna-
tional) were purchased. Rabbit and guinea pig antibodies to Ln α4, α5, and β2 are described elsewhere (Miner et al., 1997). A pAb to purified human Ln-8 (Fig. 5 B) and mAb 1G5 raised to an α4 LG1-domain fusion protein were generated in Lama40/0 mice; each labels Ln-8 on blots (Fig.

5 a) and all α4-rich BLs in normal mice, but no BLs in Lama40/0. Rabbit antibodies to integrin β1 and agrin were gifts from Eva Engvall (Barn-
ham Institute, La Jolla, CA) and David Glass (corporate license from Regeneron Pharmaceuticals), respectively. Other purchased antibodies: MAB1946 (enactin, MAB1948 [perlecan], AB1920 [integrin α3], and MAB1982 [integrin α6] from CHEMICON International; CoH3 [integrin α6; Beckman Coulter], CD29 [integrin β1; BD Biosciences]; VIA41 [α-dys-
troglycan; Upstate Biotechnology]; β-dystroglycan and β-sarcoglycan (No-
voCastra); neurofilament [2H3; Developmental Studies Hybrida Bank re-
pository, Ames, Iowa]; S100 (Neomarkers); Ki67 (Vector Laboratories); S100 (Neomarkers); Ki67 (Vector Laboratories); and S100 (Neomarkers).

Lns
Mouse Ln-1 and human Ln-2 were from CHEMICON International. Human Ln-8 was purified from T98G cell–conditioned media using nondenaturing methods. Protein precipitated by 40% ammonium sulfate was bound to a DEA- Sepharose Fast Flow FPLC column (Amersham Biosciences) in imida-
sule (10 mM; pH 7.0) and eluted with a 0.05–1.0-M NaCl gradient. Pooled fractions containing Ln-8 (0.4–0.5 M NaCl) by dot-blot assay were dialyzed (5 mM phosphate and 50 mM NaCl, pH 7.3) and repurified by DEA-Sepharose. The second DEA pool was concentrated (Aquadice; Calbiochem) and size-fractionated by FPLC through Superose 6 HR. The fi-
nal pool contained a single major protein complex at 680 kDa on silver-
stained SDS-PAGE nonreducing gels (Fig. 5 A). In this material, immuno-
activity for Ln α4, b1, and γ1 chains comigrated, and α1, α2, α5, and β2 were not detectable.

Histology
For resin sections, killed animals were perfused with 3% (wt/vol) PFA, 1% (vol/vol) glutaraldehyde, in PBS; nerves were incubated overnight at 4°C.
in 4% PFA, 4% glutaraldehyde in 0.1 M cacodylate; 1-mm pieces were
post-fixed 1 h in 1% OsO4, dehydrated through ethanol, and embedded
in Epon. Semithin sections (0.5 μm) were stained with toluidine blue (1%
in alcohol) and imaged by digital color photomicroscopy. Ultrathin sec-
tions (90 nm) stained with uranyl acetate were imaged by transmission
EM. Quantitation of myelination patterns was performed on photographic
montages of transverse sections of the peripheral nerve. Myelinating nerve
fibers were counted from semithin sections; nonmyelinated axons were
counted from ultrathin sections on Formvar-coated hole grids photo-
graphed at 2,000–10,000×.

Immunohistochemistry was done as described previously (Miner et
al., 1997), using 8–10-μm cryostat sections cut from OCT-embedded un-
fixed nerves prepared by snap freezing in −150°C 2-ethylbutyltane, or teased
sciatric nerves prepared by gently spreading 1–2-mm segments on subbed
slides. In α4 and β2 epitopes required denaturation (Miner et al., 1997).
In brief, sections were incubated overnight with antibodies di-
luted in PBS containing 5% (wt/vol) BSA, washed in PBS, and bound an-
tibodies detected with species-specific, fluorescent secondary antibodies
(1 h). Teased fibers were processed for 15 min with 2% PFA, cleared
with 0.1 M glycine, and stained in PBS with 0.5% BSA and 0.5% Triton X-100.
Hocheist 33258 (Molecular Probes, Inc.) was added to mounting medium
to visualize nuclei. TUNEL staining was performed according to kit direc-
tions (Roche; product 1684795). Myelin was visible by intrinsic fluores-
cence [Ex365 nm/Em450 nm]. Images were made at ambient tempera-
ture with PlanApo 60× [1.4 NA] oil-immersion lenses on BX microscopes
(Olympus), using either a DC500 camera and IM50 digital im-
ique ware (Leica) or an FV300 confocal scan head (Olympus). Multiply
stained images were colorized and superimposed in Photoshop 6.0.
Quantitation of Schwann cell nuclei was performed strictly on transverse
sections of PFA-fixed sciatic nerves frozen in situ (the thigh). All
nervi nuclei (Hocheist, total; Ki67, proliferating; TUNEL, apoptotic) not re-
siding in perineurial sheaths (distinguished by integrin α6 counterstaining
and morphology) were counted in digital images taken at 400× without
background subtraction.

Cell culture

Proliferation assays used Schwann cells (92–98% S100-positive) freshly
prepared from desheathed E13 chick sciatic nerves (Patton et al., 1998).
Plastic 96-wells were coated with poly-L-lysine (0.1 mg/ml, 1 h; Sigma-
Aldrich), then Ln-1, -2, or -8 in PBS (for 8 h at 4
C) and fed by 50% medium replacement at d 3, cells remained adherent to all substrates at d 5, when TUNEL assay (Roche) labeled
<2% of cells. Adhesion assays used Schwann cells cultured 10 d or less
after preparation from P4 mice. Schwann cells were enriched (>97%
S100-reactive) by complement-mediated lysis of fibroblasts with anti-
Thy-1.1 antibody (TN-26; Sigma-Aldrich), and expanded in DME, 10% FCS,
with 0.1% hyaluronidase. Cells were differentiated in 10 μg/ml retinoic
acid (RA) and incubated 3 h at 37°C. After washing with PBS, cell counts
were performed by mail order (FACS) and duplicate hemocyt-
ometer readings. Experiments included triplicate wells for each condition.

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