Both Laminin and Schwann Cell Dystroglycan Are Necessary for Proper Clustering of Sodium Channels at Nodes of Ranvier

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Nodes of Ranvier are specialized axonal domains, at which voltage-gated sodium channels cluster. How axons cluster molecules in discrete domains is mostly unknown. Both axons and glia probably provide constraining mechanisms that contribute to domain formation. Proper sodium channel clustering in peripheral nerves depends on contact from Schwann cell microvilli, where at least one molecule, gliomedin, binds the sodium channel complex and induces its clustering. Furthermore, mice lacking Schwann cell dystroglycan have aberrant microvilli and poorly clustered sodium channels. Dystroglycan could interact at the basal lamina or at the axon–glial surface. Because dystroglycan is a laminin receptor, and laminin 2 mutations [merosin-deficient congenital muscular dystrophy (MDC1A)] cause reduced nerve conduction velocity, we asked whether laminins are involved. Here, we show that the composition of both laminins and the dystroglycan complex at nodes differs from that of internodes. Mice defective in laminin 2 have poorly formed microvilli and abnormal sodium clusters. These abnormalities are similar, albeit less severe, than those of mice lacking dystroglycan. However, mice lacking all Schwann cell laminins show severe nodal abnormalities, suggesting that other laminins compensate for the lack of laminin 2. Thus, although laminins are located at a distance from the axoglial junction, they are required for proper clustering of sodium channels. Laminins, through their specific nodal receptors and cytoskeletal linkages, may participate in the formation of mechanisms that constrain clusters at nodes. Finally, abnormal sodium channel clusters are present in a patient with MDC1A, providing a molecular basis for the reduced nerve conduction velocity in this disorder.

Key words: laminin; Schwann cell; node of Ranvier; microvilli; dystrophic; sodium channels

Introduction

Nodes of Ranvier contain clusters of voltage-gated sodium channels (Nav), which are essential for saltatory nerve conduction. Laterally, paranodes and juxtaparanodes are enriched in different molecules, making the axolemma polarized (for review, see Poliak and Peles, 2003; Salzer, 2003; Scherer et al., 2004). How local axonal specialization is achieved is mostly unknown. Much evidence indicates that assembly of axonal domains depends on axoglial interactions. Molecule(s) on Schwann cell microvilli bind the nascent Ng-CAM-related cell adhesion molecule (NrCAM)/Neurofascin 186/Nav complex on axons (Vabnick et al., 1996; Lambert et al., 1997; Ching et al., 1999; Lustig et al., 2001), and one such molecule, gliomedin, has been identified recently (Eshed et al., 2005). Furthermore, axonal and glial scaffolds may form molecular barriers that prevent lateral movements (Rosenbluth, 1981; Vabnick et al., 1996; Pedraza et al., 2001). These mechanisms may induce nascent binary Nav clusters and force them to fuse, possibly pushed by elongating Schwann cell processes (Melendez-Vasquez et al., 2001; Gatto et al., 2003). However, it is hard to reconcile these data with the observation that Nav clusters form in the absence of Schwann cell processes in the roots of laminin 2-deficient mice (Deerinck et al., 1997).

Microvilli contain dystroglycan, and its ablation in Schwann cells impairs Nav clustering and reduces nerve conduction velocity (Saito et al., 2003). How dystroglycan promotes clustering is...
not known. Dystroglycan is a receptor (Ervasti and Campbell, 1993) that connects laminin and agrin in the basal lamina of internodal Schwann cells to an intracellular complex that includes dystrophin isoform 116 (Dp116), dystrophin-related protein 2 (DRP2), periaxin, and actin (Byers et al., 1993; Saito et al., 1999; Sherman et al., 2001). Similarly, dystroglycan may connect the basal lamina to the cytoskeleton of microvilli, forming the glial side of a specific molecular scaffold that restricts the movement of Nav. In this case, a distinct composition of dystroglycan linkage at the node would be expected. Alternatively, because dystroglycan can bind ligands other than laminins, dystroglycan could be on the axoglial interface of microvilli and interact with glial or axonal molecules.

To explore this question, we analyzed the composition of laminins and their linkages at nodes, and compared the phenotype of mice lacking laminins to that of mice lacking dystroglycan in Schwann cells. We show that laminins are abundant in the nodal basal laminas but that, here, the composition of both laminins and of the dystroglycan complex differs from that of internodal nodes. Mice lacking functional laminin 2 (dystrophic) have reduced nerve conduction velocity, enlarged nodes, and poorly formed microvilli, in agreement with previous data and similar to merosin-deficient congenital muscular dystrophy (MDC1A) patients (Shorer et al., 1995; Di Muzio et al., 2003). Similar Nav clusters are found in an MDC1A patient. These alterations are similar, but less severe, than those of dystroglycan-null mice. However, mice lacking all Schwann cell laminins show dystrophic microvilli, in agreement with previous data and similar to MDC1A patients (Deerinck et al., 1997), but highly abnormal. These data suggest that specific Schwann cell laminins, receptors, and cytoskeletal linkage participate in the formation of a transcellular scaffold important for the assembly of the nodal architecture and provide a molecular basis for the reduced nerve conduction velocity in patients with MDC1A.

### Table 1. List of antibodies used

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Antibody</th>
<th>Species</th>
<th>Clone/name</th>
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<th>Source</th>
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<td>DRP3/20C5</td>
<td>1:50</td>
<td>Novocastra Laboratories</td>
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mAb, Monoclonal antibody; pAb, polyclonal antibody.

### Table 2. Laminin trimer composition in adult peripheral nerves

<table>
<thead>
<tr>
<th>Laminin</th>
<th>Subunit composition</th>
<th>Perineurium</th>
<th>Endoneurium (Schwann cell basal lamina)</th>
<th>Node of Ranvier</th>
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<tr>
<td>1</td>
<td>α1β1γ1</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>2</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>α4β1γ1</td>
<td>+</td>
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<td>4</td>
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<td>+</td>
<td></td>
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<tr>
<td>6</td>
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<td>9</td>
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</tr>
<tr>
<td>10</td>
<td>α5β2γ2</td>
<td>+</td>
<td></td>
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</table>

Trimer distribution is assumed from coincident expression of individual isoforms.

### Materials and Methods

**Mice and genotyping.** *Dystrophic dy2J* mice (C57BL/6J-Lama2dy.2-1) were obtained from the The Jackson Laboratory (Bar Harbor, ME) and maintained by backcrosses with C57BL6/N. Both wild-type (wt) and dy2J+/+ were used as control for dy2J/dy2J mice. Mice lacking dystroglycan specifically in Schwann cells were produced by crossing P0Cre transgenic mice (mPoTOTTcre) (Feltri et al., 2002), with dystroglycan–floxed mice (Moore et al., 2002). DG Flox/Flox/P0Cre mice were used for the experiments, and DG+/+///P0Cre or DG Flox/Flox littermates were used as controls. Similarly, mice lacking laminin γ1 in Schwann cells were generated by crossing γ1 floxed mice (Chen and Strickland, 2003) with P0Cre mice to obtain experimental γ1 Flox/Flox/P0Cre mice and γ1 Flox/Flox control littermates. Mouse genotyping was performed by PCR analysis of tail genomic DNA (Stephens et al., 1993; Feltri et al., 1999a,b; Chen and Strickland, 2003; Saito et al., 2003). All experiments involving animals were performed according to protocols approved by the Institutional Animal Care and Use Committee of San Raffaele Scientific Institute.

**Human biopsy.** Sural nerve biopsies from a MDC1A patient and from a patient with an unrelated, acquired inflammatory neuropathy were obtained as described previously (Quattrini et al., 1996; Di Muzio et al., 2003), in accordance with protocols approved by institutional ethical committees. Immunohistochemistry on the MDC1A nerve showed absence of laminin 2 (Di Muzio et al., 2003).

**Antibodies.** The primary antibodies used are listed in Table 1. Secondary antibodies included the following: fluorescein isothiocyanate

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Immunohistochemistry on teased fibers and on frozen sections. Sciatic nerves from adult mutant and control mice and from rats; dorsal and ventral roots of dy2J/dy2J and control mice; and optic nerves from control mice at postnatal day 11 were dissected and fixed by immersion in fresh ice-cold 4% paraformaldehyde in PBS for 30 min, and then kept in PBS at +4°C until teasing. For immunofluorescence on unfixed tissues, sciatic nerves were washed in PBS after dissection and teased immediately. For quantitative analysis of Nav, mutant and control nerves were dissected and teased the same day. Single fibers from roots and sciatic nerves were teased apart after removal of the perineurium using fine stainless-steel needles, dried overnight, and stored at −20°C. For immunostaining, fibers were permeabilized in cold acetone for 10 min at −20°C. After blocking with 5% fish skin gelatin, 0.5% Triton X-100, 0.02% sodium azide in PBS, fibers were labeled with primary antibodies, followed by appropriate secondary antibodies, washed, dried, and mounted with Vectashield (Vector Laboratories, Burlington, CA), and analyzed with a confocal microscope (MRC 1024; Bio-Rad, Hercules, CA; or TCS-S2; Leica, Nussloch, Germany). Because the background fluorescence of mouse monoclonal antibodies on mouse teased fibers can resemble basa lamina staining, we used rat or rabbit antibody for laminin stainings, or rat teased fibers for dystroglycan, Dp116, and utrophin staining, in cases in which only mouse antibodies were available. Immunohistochemistry on frozen sections was performed as described previously (Feltri et al., 2002).

Electron microscopy. The sciatic nerves of wild-type and dy2J/dy2J adult mice were removed and fixed overnight at 4°C in 2% glutaraldehyde in 0.12 M phosphate buffer. On the next day, the sciatic nerves were washed in phosphate buffer, osmicated in 1% OsO4 for 1 h at room temperature, and then divided into 5-mm-long segments, dehydrated in graded ethanols, infiltrated with propylene oxide and then Epon, and polymerized at 60°C. Ultrathin sections were stained with lead citrate and photographed with a Zeiss (Oberkochen, Germany) EM10 electron microscope. Semithin morphological analysis of roots and sciatic nerves of dystrophic animals and control littermates was performed as described previously (Wrabetz et al., 2000).

Electrophysiology. Seven dy2J/dy2J and three control mice were analyzed at 3 months of age. Mice were anesthetized with tribromoethanol (0.02 ml/g of body weight) and placed under a heating lamp to avoid hypothermia. The sciatic nerve motor conduction velocity was obtained with steel monopolar needle electrodes: a pair of stimulating electrodes was inserted subcutaneously near the nerve at the ankle; and a second pair of electrodes was placed at the sciatic notch, to obtain two distinct sites of stimulation, proximal and distal, along the nerve. The compound motor action potential was recorded with an active electrode inserted in muscles in the middle of the paw and a reference needle in the skin between the first and second digits.

Image and quantitative analysis. Figures were prepared using Adobe Photoshop 7, Adobe Illustrator 10, and ImageJ. To quantify the frequency of abnormal clusters, mutant and control nerves were analyzed blindly by two independent investigators using a Leica DMS5000 fluorescence microscope: only paranodes belonging to straight fibers and in an optimal focal plane (as identified by Caspr staining of both adjacent paranodes) were chosen, and then the Nav cluster staining at the node was classified as normal (rectangular, with square corners, normal width) or abnormal (irregular shape and lacking square corners, diffuse and long, low intensity in every optical plane, or difficult to put into focus). Quantification of the are of Nav clusters in sciatic nerves and roots was made using a laser inverted confocal microscope (Leica TCS-S2) (FITC, excitation of laser Ar/Kr at 488 nm; TRITC, excitation of laser He/Ne at 546 nm). Nodes were selected using Caspr staining, as above. The gain of Nav and Caspr fluorescence detection was maintained below the threshold of fluorochrome saturation. z-axis-series spanning 5.5 μm were acquired by sequentially scanning (between frames), using a step size of 0.122 μm. ImageJ was used to measure the area of the clusters and the diameter of the axonal fibers on maximum projections of the Z series. A scale of values and a unit of length were set, and the two channels, FITC and TRITC, were split from each projection. The area of each Nav cluster to be measured was traced two to three times by two independent investigators, visually defining the perimeter as a qualitatively reduced intensity, and then calculated and averaged. Thus, this technique does not include the area of “dispersion” with low Nav staining. The average area was normalized for the average diameter of the fiber, measured by drawing a line through Caspr staining at the border between paranode and node. The values were compared by Student’s t test. The quantitative analysis of the dystrophic roots was made without normalization because of the absence of Caspr clusters.

Results
Laminins 2 and 10 are abundant in the basal lamina surrounding nodes of Ranvier
A continuous basal lamina surrounds the outer surface of Schwann cells along internodes, extending over paranodes and...
Specific members of the dystrophin–glycoprotein complex are present at nodes. Teased fibers of adult rat sciatic nerves stained with an antibody against the paranodal marker Caspr (green in B, E, H, K; red in N). The picture shows the localization of α-dystroglycan (A, arrow) and β-dystroglycan (D, arrow) at the nodes of Ranvier. The monoclonal antibody against the dystrophin isoform, Dp116, shows a striking microvillar enrichment (G, arrow). Utrophin is also present at nodes (J, arrow), whereas dystrophin-related protein 2 and periaxin are absent at nodes (M; data not shown). The merged confocal images are shown at the right (C, F, I, L, O). DG, Dystroglycan. Scale bar, 16 μm.

Dystrophic dy^2^/dy^2^ mice have minor myelin abnormalities in peripheral nerves, but reduced nerve conduction velocity

To explore whether dystroglycan alterations at nodes of Ranvier represent a loss of laminin receptor function, we compared the phenotypes of dystroglycan-null mice and dy^2^/dy^2^ mice. Older studies showed that dystrophic mice have reduced nerve conduction velocity (Rasminsky et al., 1978) and widened nodes of Ranvier (Bradley et al., 1977; Jaros and Bradley, 1979; Jaros and Jenkinson, 1983). These studies were performed on dy/dy mice, which lack laminin 2 because of an unidentified genetic defect mapping to the laminin α2 locus (Sunada et al., 1994). To compare the phenotype of laminin 2 mutants to that of dystroglycan mutants, we used dystrophic dy^2^/dy^2^ mice instead, which have nonfunctional laminin 2 because of a known mutation in the α2 chain. First, we validated the model by showing that, similarly to dy/dy mice, dystrophic dy^2^/dy^2^ mice have a 30% reduction in nerve conduction velocity and an increased F-wave latency (Fig. 3). Dystrophic dy/dy and dy^2^/dy^2^ mice have defective radial sorting of axons that results in bundles of naked axons, more severe in spinal roots than in distal nerves (Bradley and Jenkinson, 1973). To evaluate the potential contribution of the distal sorting defects to the observed reduction in nerve conduction velocity, we compared semithin sections of roots and sciatic nerves in dy^2^/dy^2^ mice. As shown in Figure 3, the radial sorting defect was much more pronounced in roots than in distal nerves (Fig. 3) (Bradley and Jenkinson, 1975; Weinberg et al., 1975). Thus, axonal sorting defects in nerves do not likely explain the observed reduction in nerve conduction velocity in the nerves themselves. Therefore, mice lacking laminin 2 phenocopy the slowing in nerve conduction velocity seen in mice lacking Schwann cell dystroglycan (Saito et al., 2003).
Dystrophic \(dy^{2J}/dy^{2J}\) mice have abnormal nodes of Ranvier and microvilli

To study microvilli in \(dy^{2J}/dy^{2J}\) mice, we examined the ultrastructure of nodes in longitudinal sections of sciatic nerves from \(dy^{2J}/dy^{2J}\) mice and control nerves. As shown in Figure 4, some nodes in dystrophic nerves were abnormally wide (>5 μm), with poorly formed microvilli. These abnormalities were present at various degrees in ~20% of nodes. In extreme cases, microvilli were absent, such as the node shown in Figure 4A.

Next, we determined whether the molecular components of the nodal region were altered in \(dy^{2J}/dy^{2J}\). Staining with microvilli markers showed normal staining for Dp116 in \(dy^{2J}/dy^{2J}\) mice (Fig. 4) as well as for radixin, moesin, and ezrin (Scherer et al., 2001) in the majority of \(dy^{2J}/dy^{2J}\) and dystroglycan-null nodes (Fig. 4 and data not shown). ERM protein phosphorylation, as determined by staining with a phosphospecific antibody, appeared normal in many nodes, but was reduced or absent in some nodes of both dystrophic and dystroglycan-null nerves (Fig. 4) (data not shown). To analyze the localization of paranodal and juxtaparanodal molecules in \(dy^{2J}/dy^{2J}\) mice, we stained teased sciatic nerve fibers with antibodies recognizing paranodal (Caspr on axons and Neurofascin 155 on Schwann cells) and juxtaparanodal (Kv1.1 potassium channels) markers. Supplemental Figure 1 (available at www.jneurosci.org as supplemental material) shows that, similarly to mice lacking Schwann cell dystroglycan, glial and axonal paranodal and juxtaparanodal markers are correctly localized in \(dy^{2J}/dy^{2J}\) nerves. Thus, nodes lacking functional laminin 2 have various degrees of abnormalities that phenocopy those seen in mice lacking Schwann cell dystroglycan.

Dystrophic \(dy^{2J}/dy^{2J}\) mice have abnormal Nav channel clusters

To determine whether Nav clusters were also perturbed in \(dy^{2J}/dy^{2J}\) mice, we compared Nav staining on teased sciatic nerves from \(dy^{2J}/dy^{2J}\) mice to that in control nerves and in nerves lacking Schwann cell dystroglycan. Antibodies directed against all Nav isoforms, or to Nav1.6, the major isoform present at mature nodes (Boiko et al., 2001), were used. In nerves from \(dy^{2J}/dy^{2J}\) animals, Nav clusters were often smaller, stained with lower intensity, and were more irregularly shaped than those of control animals. Whereas clusters in wild-type animals were rectangular and had square corners, \(dy^{2J}/dy^{2J}\) clusters were irregular and lacked definite corners (Fig. 5, compare A with G and I–K with R–T). Some clusters were dispersed laterally (Fig. 5I). Because paranodal junctions were normal (data not shown) it is possible that laterally diffused clusters are present in abnormally wide nodes of Ranvier. “Binary nodes,” two distinct Nav clusters on the same axon separated by space, were rarely seen. These abnormalities were identical in quality but less severe than those observed in nerves lacking dystroglycan (Saito et al., 2003), in which the low intensity and lateral dispersion of Nav were more extreme (Fig. 4 and data not shown). ERM protein phosphorylation, as determined by staining with a phosphospecific antibody, appeared normal in many nodes, but was reduced or absent in some nodes of both dystrophic and dystroglycan-null nerves (Fig. 4) (data not shown). To analyze the localization of paranodal and juxtaparanodal molecules in \(dy^{2J}/dy^{2J}\) mice, we stained teased sciatic nerve fibers with antibodies recognizing paranodal (Caspr on axons and Neurofascin 155 on Schwann cells) and juxtaparanodal (Kv1.1 potassium channels) markers. Supplemental Figure 1 (available at www.jneurosci.org as supplemental material) shows that, similarly to mice lacking Schwann cell dystroglycan, glial and axonal paranodal and juxtaparanodal markers are correctly localized in \(dy^{2J}/dy^{2J}\) nerves. Thus, nodes lacking functional laminin 2 have various degrees of abnormalities that phenocopy those seen in mice lacking Schwann cell dystroglycan.

Nerves of mice lacking all Schwann cell laminins have widespread nodal abnormalities

The above data show that nodal abnormalities were more severe in mice lacking dystroglycan than in mice lacking laminin 2. To
evaluate whether this was attributable to redundant/compensatory function of other laminins present at nodes of Ranvier of dystrophic mice (laminin 1 and laminin 10), we studied Nav clustering in mice in which the laminin 1 chain gene, common to all of the laminin isoforms present in Schwann cells, was specifically ablated in Schwann cells using the Cre/loxP system (P0CreLAMA2null). In these nerves, most axons are naked because of defective radial sorting (Yu et al., 2005). In the few myelinated fibers, recognizable by paranodal Caspr staining, Nav clusters presented the same abnormalities described in dystrophic mice, but more severe (Fig. 5 E, O–Q) and at higher frequency (59% of abnormal clusters; \( p = 0.001 \) by \( \chi^2 \) test when compared with controls). Thus, compensation by laminin 1 and/or redundancy with laminin 10 (Yang et al., 2005) probably partially substitute for the lack of laminin 2 in dystrophic nodes.

Severe Nav cluster abnormalities are present in roots of \( dy^{2J}/dy^{2J} \) mice

The presence of Nav clusters in naked axons of dystrophic mice roots have been reported (Deerinck et al., 1997) and cited as evidence that microvilli and glial contact are not strictly necessary for Nav clustering. Having classified normal and abnormal Nav clusters in mutant dystrophic and dystroglycan mice, we asked whether clusters on dystrophic roots were normal. Dystrophic roots contained naked axons and myelinated fibers (Fig. 3), distinguishable by the absence or presence of Caspr paranodal staining, respectively. Indeed, Nav clusters could be seen in amyelinated areas, but they were extremely small, faintly stained, and irregularly shaped or diffuse (Fig. 6). Comparison between naked and myelinated axons showed that the abnormalities were clearly most severe in amylinated axons (Fig. 6). Microvilli were absent at these sites, as shown by lack of radixin staining (Fig. 6). Thus, in the absence of glial processes, Nav clusters are recognizable, but highly abnormal.

The area and density of Nav clusters are reduced in the absence of laminins and dystroglycan

So far, we presented evidence that Nav clusters are qualitatively abnormal when Schwann cells lack laminins or dystroglycan, and these abnormalities correlate with abnormalities in microvilli at nodes of Ranvier. Furthermore, when microvilli are completely missing, Nav clusters appear most abnormal. To quantify these differences, we measured the size of Nav clusters, stained with anti-Nav antibodies, in nerves from mice lacking laminin 2 (\( dy^{2J}/dy^{2J} \)), all laminins (P0Cre\( \gamma \)) or dystroglycan (P0Cre DG), and in naked roots lacking microvilli completely (\( dy^{2J}/dy^{2J} \)). As shown in Figure 7, the area occupied by Nav clusters was similarly reduced in mice lacking laminins or dystroglycan, and more severely reduced in naked axons from dystrophic roots. Thus, the severity in reduction of Nav clusters area correlates with the severity of microvilli abnormalities. Reduction of the area of Nav clusters was not attributable to a reduction in fiber diameters in mutant nerves, because it remained smaller than in control nerves when corrected for fiber diameter (as estimated by Caspr staining) (Fig. 7).

Abnormal Nav clusters in a MDC1A patient

\( dy^{2J}/dy^{2J} \) mice are a model for MDC1A. MDC1A patients are heterogeneous, but most have reduced nerve conduction velocities (Shorer et al., 1995; Quijano-Roy et al., 2004). To determine whether abnormalities in Nav clustering can account for slowing of conduction velocity, we evaluated an MDC1A patient with a Gly600Arg mutation in the LAMA2 gene, in which a peripheral neuropathy with slow nerve conduction has been described previously (Di Muzio et al., 2003). Double staining of frozen sections from the sural biopsy of the MDC1A patient with anti-Nav and anti-Caspr antibodies showed abnormalities in Nav clusters (12 of 18 abnormal nodes), when compared with those of a patient with an unrelated inflammatory neuropathy (7 of 23 abnormal nodes) (\( p < 0.02 \) by \( \chi^2 \) test) (Fig. 8). Thus, the laminin 2 deficit in humans causes Nav clustering abnormalities similar to those observed in mouse models lacking laminins or dystroglycan. These abnormalities likely contribute to the observed reduction in nerve conduction velocity.

Discussion

In this paper, we show that the node of Ranvier contains a specific repertoire of laminins and dystrophin linkages to the cytoskele-
ton. Thus, the discrete microdomains of myelinated axons also extend to the external Schwann cell surface. In addition, we show that eliminating one or more components of these surface microdomains is sufficient to cause nodal abnormalities, including abnormal microvilli and abnormal clusters of Nav, which likely contribute to reduced conduction velocities in mice and in MDC1A patients.

Specific laminins and specific dystrophin—glycoprotein complexes at nodes

Myelinated axons and the overlying Schwann cells are longitudinally polarized in corresponding domains (Poliak and Pelles, 2003; Salzer, 2003). We find that the basal lamina, a continuous structure outside the Schwann cells (as is the axon inside), is also longitudinally polarized, because the nodal—paranodal—juxtaparanodal region has a different laminin composition than the rest of the internode. In addition, this regional difference appears to be accompanied by a different cytoskeletal linkage through different dystroglycan complexes. This resembles the situation in muscle fibers and in other tissues, in which laminin, laminin receptors, and cytoskeletal components are organized in discrete domains, influencing cell polarization and tissue morphogenesis (for review, see Yurchenco et al., 2004). Thus, the entire basal lamina/Schwann cell/axon unit is polarized, and laminins may play an essential role in the polarization and morphogenesis of the peripheral nerve unit.

Laminins cooperate with dystroglycan at nodes

We find that the nodal abnormalities of laminin-deficient Schwann cells resemble those of dystroglycan-deficient Schwann cells. In the absence of laminin 2, Schwann cells have poorly formed and misoriented microvilli, altered Nav clustering that is qualitatively similar to dystroglycan-null Schwann cells, and normal paranodes and juxtaparanodes. The quantitative differences in severity between the two mutants can be explained by redundancy with laminin 10 or compensation by laminin 1 in dystrophic mice. That the phenotypes overlap at the node of Ranvier strongly suggests that laminins and dystroglycan are in the same genetic pathway that promotes nodal Nav clustering. Because laminin is a ligand for dystroglycan in Schwann cells (Yamada et al., 1996), it is plausible that there is an epistatic relationship between them.

Laminins and dystroglycan could promote Nav clustering in two, non-mutually exclusive, ways. First, the binding of dystroglycan to laminin transmits signals that organize the cytoskeleton of microvilli, contributing to mechanisms that form or constrain Nav clusters. This interaction need not occur only at nodes, and could maintain proper orientation of microvilli toward the axon so that molecules on microvilli tips, such as gliomedin, would be allowed to bind the NrCAM/Neurofascin 186/Nav complex. This is a conservative interpretation of our data and fits with the recent finding that dystroglycan regulates the actin cytoskeleton through ezrin (Spence et al., 2004). However, at least at the level of immunohistochemistry, ERM, ERM phosphorylation, and Dp116 were not severely disrupted in mutant mice.

Second, dystroglycan could have a more direct role in microvilli, distinct from laminin binding, by binding other microvillar or axonal components of the node. Gliomedin has been shown recently to bind the NrCAM/Neurofascin 186/Nav complex and to enculture Nav clusters on cultured DRG axons (Eshed et al., 2005). Gliomedin contains a collagen motif that could interact with molecules such as dystroglycan on microvilli (Eshed et al., 2005). Alternatively, Schwann cell dystroglycan can bind agrin (Yamada et al., 1996), and in other cell types dystroglycan can bind perlecan, biglycan, and neurexin (Talts et al., 1999; Bowe et al., 2000; Sugita et al., 2001). The role of non-neuronal agrin in peripheral nerve is unknown, but Schwann cells contain active agrin isoforms with acetylcholine receptor clusterizing activity (Yang et al., 2001). Thus, dystroglycan could immobilize diffusible agrin in a proteoglycan complex to facilitate Nav clustering, as it happens for acetylcholine receptors at neuromuscular junctions (Grady et al., 2000; Jacobson et al., 2001). In these interpretations, laminin would bind a different receptor over nodes (e.g., α6β1 integrin), and dystroglycan would have a different binding partner (e.g., agrin, proteoglycans, Neurofascin 186) and more directly cluster or bind Nav on the axonal membrane.

Laminins and their actin linkage may participate in the formation of barriers or molecular sieves at the node

The formation of the axoglial apparatus at nodes of Ranvier probably depends on a temporal hierarchy of sorting, targeting, as-
Among the constraining mechanisms, molecular sieves or barriers have been proposed that could restrict the lateral passage of Nav, maintaining their selective localization at nodes (Rosenbluth, 1981; Vabnick et al., 1996). These sieves comprise a molecular scaffold that is linked to the cytoskeleton by transmembrane proteins, that spans both the axon and the Schwann cell, and that depends on trans interaction at the axoglial junction. Finally these molecular scaffolds could operate during development and constraining mechanisms (Pedraza et al., 2001).

Figure 6. Severe abnormalities of Nav clusters in naked axons of dystrophic roots. Confocal images of Nav clusters in dystrophic (A, B, E, F) and normal (C, D, G, H) teased spinal roots immunolabeled with anti-Nav1.6 (green) and anti-Caspr (red) antibodies in A–D, or anti-radixin (green) and anti-panNav (red) in E–H. Faint Nav immunoreactivity is present at discrete sites along dystrophic fibers (A, arrows and enlargement), likely corresponding to naked axons because of a lack of the paranodal Caspr (red) and nuclear 4′,6′-diamidino-2-phenylindole (DAPI) (blue) markers (B). Nav clusters found in these regions are small, irregularly shaped, and diffusive along the fiber (A, B, arrows), compared with Nav clusters from wild-type spinal roots (C, D). These clusters form in the absence of microvilli, as shown by the lack of radixin staining (E, F, and enlargements). Scale bar, 16 μm.

Figure 7. Decrease in the area of Nav channel clusters in the absence of laminins or dystroglycan. Quantitative analysis of the area occupied by Nav clusters in nodes from dystrophic, laminin γ1-null, and dystroglycan-null sciatic nerves and dystrophic roots. Open columns represent the average areas stained with anti-Nav1.6 antibodies. Filled columns show the same data after normalization for fiber diameter. A reduction in the area of Nav clusters is evident in dystrophic dy2/dy2, laminin γ1, and dystroglycan-null sciatic nerves compared with controls. Similar quantitative analysis in dystrophic roots shows a severe reduction in the area of Nav clusters compared with wild-type roots.

Figure 8. Nav clusters abnormalities in an MDC1A patient. Longitudinal sections of sural nerve biopsies from an MDC1A patient (A, B) and a control sural nerve (C, D) double stained with an antibody that recognizes all isoforms of sodium channels (PanNa+, red) and an anti-Caspr antibody (green). A, B, The image shows a reduction in the intensity of staining for Nav clusters (arrows, enlarged in insets) on MDC1A nerves compared with control nerves. Scale bar, 16 μm.

Table 1. Summary of the average Nav channel cluster area and average diameter in sciatic nerves and roots.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Average Nav cluster area</th>
<th>Average diameter</th>
<th>Student t-test mutant/wt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type (n=23)</td>
<td>2.480</td>
<td>1.521</td>
<td></td>
</tr>
<tr>
<td>dy2/dy2 (n=50)</td>
<td>1.354</td>
<td>1.366</td>
<td>p&lt;0.0001</td>
</tr>
<tr>
<td>POCre γ1 (n=21)</td>
<td>1.313</td>
<td>1.297</td>
<td>p&lt;0.0001</td>
</tr>
<tr>
<td>POCre DG (n=58)</td>
<td>1.249</td>
<td>1.316</td>
<td>p&gt;0.0001</td>
</tr>
<tr>
<td>Wild type roots</td>
<td>1.932</td>
<td>1.925</td>
<td></td>
</tr>
<tr>
<td>dy2/dy2 roots</td>
<td>0.861</td>
<td>0.891</td>
<td>p&lt;0.0001</td>
</tr>
</tbody>
</table>
ment, by “pushing” Nav from internodes toward the node, driven by the longitudinal elongation of Schwann cells (Dugandzija-Novakovic et al., 1995; Pedraza et al., 2001). This model predicts that different domains of specific glial and axonal molecules must exist, that actin binding proteins are important in the formation of the scaffold, and that the lateral force mounted by the elongating Schwann cell during development would be crucial to properly compact Nav clusters at nodes. Strikingly, all of the data presented here fit with this model, because we show that there is a precise distribution of selective laminins and transmembrane and actin-binding proteins at the outer Schwann cell surface above the nodes of Ranvier, and that these molecules are crucial for the normal nodal architecture.

Role of microvilli in Nav clustering

We show that the most severe abnormalities in Nav clusters were present on the naked axons found in dystrophic roots and laminin γ1-null sciatric nerves. Here, Schwann cell processes, and therefore microvilli, are completely absent. These data extend the work of Deerinck et al. (1997), who noted the presence of Nav clusters in naked dystrophic roots but did not classify them as normal or abnormal. In keeping with our analysis, the dystrophic clusters presented in Figure 1 of their paper are irregularly shaped and diffuse. Thus, we propose that, in vivo, axons alone can initiate nucleation of Nav clusters, but also that microvilli are required to catalyze and/or maintain proper clusters, akin to the prepatterned versus stable synaptotrophic contact formation of acetylcholine receptor clusters at neuromuscular junctions (Burden, 2002). Interestingly, dystroglycan modulates this transition at neuromuscular junctions (Grady et al., 2000; Jacobson et al., 2001). It is important to note that a naked axon in a dystrophic mouse may not be devoid of Schwann cells throughout its full length, but may enter and exit from myelinated regions. Thus, any naked axon is in contact with Schwann cells at some point, and may receive instructive signals that are contact dependent, yet operate from the distance. An alternative possibility is that Schwann cell processes were present, and then retracted, around axons in bundles. This phenomena has been observed around axons of mice lacking Schwann cell β1 integrin (Feltri et al., 2002), but not γ1 laminin (Yu et al., 2005). In this case the role of microvilli would be not only to maintain but also to nucleate clusters. In both cases, microvilli are required for efficient formation and maintenance of Nav clusters. This function of microvilli is impaired in the absence of laminins or dystroglycan.

Role of Nav clustering in the pathogenesis of congenital muscular dystrophy neuropathy

Deficit of laminin 2 in humans and mice causes a reduction in nerve conduction velocity of ~30% (Rasminsky et al., 1978; Shorer et al., 1995; Di Muzio et al., 2003). Several factors can account for this reduction, including a reported reduction of density of myelinated axons (Montgomery and Swenarchuk, 1978), as well as a reduction in internodal length (Jaros and Jenkison, 1983; Di Muzio et al., 2003; Court et al., 2004). Here, we add another potential contributor by showing that, in addition to widened nodes of Ranvier (Bradley et al., 1977; Di Muzio et al., 2003), the Nav channel density is probably reduced in both dy/2 and an MDC1A patient. Although it would be difficult to dissect the role of each of these factors to reduce nerve conduction velocity, all of them could contribute to the pathogenesis of the MDC1A neuropathy.

References

Jaros E, Jenkison M (1983) Quantitative studies of the abnormal axon-


