Integrin-linked kinase is required for laminin-2–induced oligodendrocyte cell spreading and CNS myelination

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Early steps in myelination in the central nervous system (CNS) include a specialized and extreme form of cell spreading in which oligodendrocytes extend large lamellae that spiral around axons to form myelin. Recent studies have demonstrated that laminin-2 (LN-2; α2β1γ1) stimulates oligodendrocytes to extend elaborate membrane sheets in vitro (cell spreading), mediated by integrin α6β1. Although a congenital LN-2 deficiency in humans is associated with CNS white matter changes, LN-2–deficient (dy/dy) mice have shown abnormalities primarily within the peripheral nervous system. Here, we demonstrate a critical role for LN-2 in CNS myelination by showing that dy/dy mice have quantitative and morphologic defects in CNS myelin. We have defined the molecular pathway through which LN-2 signals oligodendrocyte cell spreading by demonstrating requirements for phosphoinositide 3-kinase activity and integrin-linked kinase (ILK). Interaction of oligodendrocytes with LN-2 stimulates ILK activity. A dominant negative ILK inhibits LN-2–induced myelin-like membrane formation. A critical component of the myelination signaling cascade includes LN-2 and integrin signals through ILK.

Introduction

Formation of myelin is a function of oligodendrocytes in the central nervous system (CNS) and Schwann cells in the peripheral nervous system (PNS) in subtle interaction with adjacent axons. To decipher the molecular mechanisms involved in this process in the CNS, we have reduced myelination to three fundamental but likely interrelated cell biological mechanisms: (1) a “cell spreading” machinery to form the large surface membrane characteristic of differentiated oligodendrocytes; (2) specific cell-cell recognition molecules that stabilize oligodendrocyte–axon interactions; and (3) a molecular motor that drives spiraling of the oligodendrocyte’s leading edge around the axon.

Here, we have focused on cell spreading, which involves the elaboration of large membrane sheets, and on the genesis of specialized sites on the cell surface for contact with axons. Clues to the molecular mechanisms involved have come from previous studies, which showed that laminin-2 (LN-2; α2β1γ1), also called merosin, promotes extensive membrane production by oligodendrocytes through integrin α6β1 in vitro (Buttery and ffrench-Constant, 1999). Importantly, LN-2 is expressed on the axonal surface in the developing CNS (Colognato et al., 2002), whereas in the developing PNS, it is expressed in the basal lamina. Furthermore, in the PNS, integrin β1–null Schwann cells migrate, proliferate and survive, but have a defect in the ensheathment of axons and formation of myelin (Feltri et al., 2002). These results prompted us to ask whether LN-2 and integrin β1 might also be important for cell spreading and myelination by CNS oligodendrocytes.

Consistent with the studies described above, congenital LN-2 deficiency in humans is associated with CNS white matter changes as shown by magnetic resonance imaging (Farina et al., 1998). However, LN-2–deficient (Lama2−/−) mice (formerly symbolized as dy, dystrophia muscularis, or mer, merosin; we refer to the affected homozygotes hereafter as, dy/dy) have been reported to exhibit primarily a neuro-
muscular phenotype including peripheral nerve hypomyelination and dysmyelination, with little evidence of defects in CNS myelination (Tsui and Matsushita, 1985; Matsumura et al., 1997). The β1 integrin signal can function through integrin-linked kinase (ILK), a 59-kD multidomain focal adhesion (FA) protein, which functions as a molecular scaffold at cell-ECM adhesion sites and participates in signal transduction pathways that control cell survival, differentiation, morphology, proliferation, and gene expression in mammalian cells (Wu and Dedhar, 2001). Within FA complexes, ILK can interact with other FA proteins such as paxillin, affixin, and CH-ILKBP (Tu et al., 2001). In Caenorhabditis elegans and Drosophila melanogaster, loss of function mutations in ILK and integrin-null mutations result in comparable phenotypes (Zervas et al., 2001; Mackinnon et al., 2002). The activation of ILK is phosphoinositide 3-kinase (PI3K) dependent, with PI 3,4,5-triphosphate, a PI3K lipid product, binding to the pleckstrin homology-like domain of ILK and activating its serine/threonine kinase activity (Dedhar, 2000).

Here, we have further defined the phenotype of dy/dy mice by identifying hypomyelination in small-diameter axons and dysmyelination in large-diameter axons within the CNS. We have determined the relevant molecular signaling steps for LN-2–induced oligodendrocyte cell spreading and myelin formation in vitro. We show that PI3K activity is required for oligodendrocytes to extend large cytoplasmic sheets, independent of effects on survival, and that ILK is localized within FA complexes of oligodendrocytes and physically associates with other FA proteins such as paxillin. In vivo, ILK is expressed primarily in oligodendrocytes within CNS white matter tracts. Finally, a dominant negative (DN) functioning ILK mutant gene blocks formation of the giant sheets of oligodendrocytes.

We conclude that dy/dy mice have defects in CNS myelination, and that the molecular events involved in oligodendrocyte cell spreading and myelin formation include signal transduction cascades using LN-2, integrin β1, PI3K, ILK, and possibly AKT.

Results

dy/dy mice are myelin-deficient and have fewer mature oligodendrocytes

A mutation (Lama2$^{△β}$) in the laminin-α2 chain gene causes of LN-2 deficiency in dy/dy mice. These mice show hindleg paralysis at ~3.5 wk. To test the hypothesis that LN-2 is involved in CNS myelin formation, we first quantified total cell concentration and total mature oligodendrocyte cell concentration in coronal hematoxylin and eosin (H&E)- and CC1-stained sections, respectively, through the corpus callosum (CC; Fig. 1 A). The CC1 antibody recognizes oligodendrocyte cell bodies without labeling myelin (Fuss et al., 2000). We observed a normal concentration of total cells and a 30% reduced concentration of CC1+ oligodendrocytes in the CC of dy/dy mice as compared with controls at matched anatomical sites (P < 0.05; Fig. 1 B). Despite the normal total cell concentration, the total number of cells was reduced, because the CC area was smaller in the mutants (Fig. 1 A), and the number of mature oligodendrocytes was reduced even more than was indicated by the measured 30% reduction in concentration. The decrease in CC1+ oligodendrocytes was accompanied by a reduction in myelin basic protein (MBP) content in the forebrain determined by immunoblot (Fig. 1 C). In contrast, MBP content in the spinal cord (SC) of dy/dy and control mice was equivalent (Fig. 1 C).
The above results suggest a defect in forebrain myelination in dy/dy mice. To determine whether myelin structure as well as myelin content is compromised, we analyzed by EM sagittal sections of CC (Fig. 2), transverse sections of optic nerve (ON; Fig. 3), and corticospinal tract in transverse sections of cervical SC (Fig. 3). In the CC, we divided myelinated axons into large diameter (axons > 0.5 μm), and small diameter (axons < 0.5 μm) groups and observed ∼30% reduction in the number of small diameter axons per unit area that were myelinated in dy/dy mice (P < 0.005; Fig. 2, A and B). The total number of axons per unit area was not changed (Table I). Higher magnification revealed that ∼40% of myelinated axons in dy/dy mice showed ultrastructural abnormalities (P < 0.0001) with large pockets of oligodendrocyte cyto-
Oligodendrocytes from dy/dy brain (CC) are more responsive to LN-2 than those from the dy/dy SC.

(A) Levels of LN-2 (M-chain, 80kD) were detected by immunoblot. The lysates were prepared from the SN and 10 d-mixed glial cell cultures from the CC area of brain and from the SC of P day 1 pups. The dy/dy mice show no LN-2 expression in the SN and reduced levels in the brain, compared with controls (n = 3). β-Tubulin levels were detected as a control. (B) The mixed glial cells from the brain and SC were cultured in 2% FBS for 10 d on poly-L-ornithine (PLO) or LN-2 and stained for O1 (green), as a marker for maturing oligodendrocytes, and DAPI (blue) for nuclei. Bar, 100 μm. 10 and 5% of cells from the brain and the SC showed O1+, respectively. The dy/dy cells on PLO generally failed to show cell spreading, in dramatic contrast to those cultured on LN-2. (C) The relative percentages of O1+ cells with broad myelin membrane sheets were counted. In confirmation of the image data in B, O1+ cells prepared from dy/dy brain on LN-2 showed dramatically increased oligodendrocyte cell spreading compared with those on PLO (P < 0.0001, asterisk) or compared with O1+ cells from control brain on PLO (P < 0.05, asterisks). Although O1+ cells prepared from the SC of dy/dy mice showed an increase on LN-2 than on PLO or control, the difference was not significant. Results are presented as ±SEM; and comparisons by ANOVA are significant (indicated by asterisks).

Table I. Concentration of small diameter myelinated axons is decreased in dy/dy mice

<table>
<thead>
<tr>
<th>Total axons/mm²</th>
<th>Myelinated axons</th>
<th>Unmyelinated axons</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[L]</td>
<td>[S]</td>
</tr>
<tr>
<td>ctrl</td>
<td>228 ± 12.4</td>
<td>40 ± 2.8</td>
</tr>
<tr>
<td>dy/dy</td>
<td>228 ± 11.1</td>
<td>40 ± 1.2</td>
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The number of large and small diameter axons that are myelinated and unmyelinated (total axons) per unit area (mm²) in the control and in the dy/dy mice was counted. t test revealed a statistically significant reduction of small myelinated axons in dy/dy mice (P < 0.05); [L] = large diameter axons (>0.5 μm); and [S] = small diameter axons (<0.5 μm).

plasm separating myelin lamellae (Fig. 2, A and C). Mean myelin sheath thickness in dy/dy mice was ~0.04 μm compared with the thickness of ~0.09 μm in controls (P < 0.0001). The average diameter of axons in control and dy/dy mice did not differ significantly; the difference in mean g-ratio (axonal diameter/total fiber diameter) of 0.82 for control fibers versus 0.91 in dy/dy mice (P < 0.0001; Fig. 2 D) indicates myelin reduction in the mutants.

In the ON from dy/dy mice, ~15% of the axons were dysmyelinated compared with none in controls (P < 0.05; Fig. 3 A and C). The total cross-sectional area of the nerve and the number of myelinated axons per unit area were not changed (Fig. 3 B), but the g-ratio was significantly increased compared with controls (P < 0.001; Fig. 3 D), to the same extent as in the CC. In the SC, consistent with the levels of MBP staining by immunoblot (Fig. 1 C), the total number of myelinated axons (Fig. 3 B), myelin morphology (Fig. 3 C) and myelin thickness (not depicted) were normal. These results highlight the regional differences in myelination in dy/dy mice.

Responsiveness of oligodendrocytes to LN-2 in dy/dy mice: regional differences

Given the regional specificity of myelination abnormalities in dy/dy mice, we next explored whether oligodendrocytes from different regions of these mice show different behavior on LN-2 in vitro. Although the mutation in dy/dy mice has not yet been identified, these mice express small amounts of apparently normal LN-2 in muscle, but no LN-2 in peripheral nerves (Guo et al., 2003). We cultured mixed glial cell preparations from the CC region of the forebrain and from the SC of postnatal (P) day 1 pups for 10 d and labeled cells with mAb to galactocerebroside (O1).

10 and 5% of cells from the brain and the SC, respectively, showed O1 immunoreactivity. The dy/dy pups were distinguished from +/+ and +/dy littermates in immunoblots by showing markedly reduced levels of LN-2 (M-chain) in the sciatic nerve (SN) and mixed glial cultures of the brain, respectively, compared with controls (Fig. 4 A). The dy/dy cells on poly-L-ornithine (PLO) generally failed to show cell spreading, in dramatic contrast to those cultured on LN-2 (Fig. 4 B). O1+ cells prepared from dy/dy brain on LN-2 showed dramatically increased oligodendrocyte cell spreading compared with those on PLO (P < 0.0001) or compared with O1+ cells from control brain on PLO (P < 0.05; Fig. 4 C). Although O1+ cells prepared from the SC of dy/dy mice showed larger processes on LN-2 than on PLO or
Buttery and ffrench-Constant (1999) in which ECM components influence oligodendrocyte morphology. We plated oligodendrocyte precursor cells (OPCs) on four different types of ECM and labeled with oligodendrocyte-specific antibodies to O1 and MBP.

O1 cells grown on LN-2 extended broad sheet-like processes in contrast to cells on BSA, thrombospondin-1 (TSP-1), or tenascin-C (TN-C; Fig. 5 A), and the difference was highly significant (P < 0.0001; Fig. 5 B). Average O1 cell diameter with sheets was 140 μm ± 10 in the LN-2–treated cells compared with 90 μm ± 15 in the other ECMs-treated cells. In addition, LN-2 enhanced the percentage of MBP+ cells (P < 0.0001) and increased the MBP levels detected by immunoblot (Fig. 5 C; unpublished data). The increase in number of O1+ cells bearing sheets was not due to increased cell survival on LN-2. Cells were plated on different ECMs for 6 h, 1 d, 2 d and 3 d and cell survival was measured through the Live/Dead Cytotoxicity and the Alamar blue assays. Both assays indicated no statistical differences in survival between the substrates (Fig. 5 D).

**LN-2 promotes oligodendrocyte cell spreading through PI3K, but not MAPK**

Because PI3K is involved in integrin β1–mediated cell spreading in various cell types, we hypothesized that LN-2 may enhance oligodendrocyte cell spreading through PI3K. PI3K activity was determined by measuring activation of AKT, a primary downstream substrate of PI3K. OPCs were replated on different ECMs, and 45 min and 6 h later the level of phosphorylation was determined by immunoblot with a phospho-AKT ser473 antibody (Fig. 6 A). LN-2, TSP-1, and TN-C all activated the phosphorylation of AKT at 45 min. However, LN-2 activated AKT phosphorylation continuously up to 6 h in contrast to other ECMs, suggesting that AKT phosphorylation might be important for LN-2–induced cell spreading in oligodendrocytes.

Because LN-2 promotes membrane production through integrin α6β1 and both PI3K and MAPK pathways are activated by integrin α6β1, we sought possible involvement of these two molecular pathways in the effects of LN-2 on oligodendrocytes in vitro. These pathways were disrupted with small molecule inhibitors: 0.01 μM wortmannin (irreversible PI3K inhibitor); 0.5 μM LY294002 (reversible PI3K inhibitor); and 4 μM U0126 (MAPK/ERK inhibitor). These inhibitors blocked the phosphorylation of AKT and MAPK in a time-dependent manner (Fig. 6, B–D). Inhibitors of PI3K but not MAPK pathways disrupted LN-2–enhanced membrane sheet formation (Fig. 6 E). 30% of O1+ cells on LN-2 formed broad myelin membrane sheets (Fig. 6 G). However, the inhibitors of PI3K blocked formation of these sheetlike processes; >50% of cells with long processes lacked membrane sheets (P < 0.005). We found no influence of MAPK inhibitor on LN-2–induced cell spreading (Fig. 6, E and G). Together, these results suggest that LN-2 promotes oligodendrocyte cell spreading through the PI3K but not the MAPK pathway.

Because AKT is one of crucial downstream effectors of PI3K, we examined the functional role of AKT in LN-2–induced oligodendrocyte cell spreading by using a DN-
AKT (kinase inactive), in which both phosphorylation sites of (Thr308 and Ser473) were replaced by alanine (AA-AKT; Kitamura et al., 1998). HA-epitope–tagged DN-AKT was expressed in the cells after incubation with tetracycline responsive promoter (TRE)-DN-AKT and tetracycline-controlled transactivator (tTA) for 3 d by adenovirus-mediated Tet–off inducible system. Although GFP vector alone (20 multiplicity of infection [MOI])-transfected O1/H11001 cells on LN-2 formed 50% broad sheetlike processes consistent with myelin membrane sheets in 3 d (Fig. 6, F and H), TRE-DN-AKT (10 MOI) and tTA (10 MOI)-transfected cells decreased LN-2-induced cell spreading by nearly 75% (Fig. 6, F and H; P < 0.0001). However, additional incubation with 10 μg/ml Tet turned off the expression of DN-AKT, showing as much cell spreading as with GFP-vector alone (Fig. 6, F and H). Furthermore, there was no statistically significant difference on cell survival between GFP-vector alone– and DN-AKT–transfected cells, but addition of Tet increased cell survival (Fig. 6 I, P < 0.5). These results show that AKT activation might be involved in LN-2–enhanced cell spreading in oligodendrocytes.

ILK is localized in oligodendrocytes in vivo and in vitro
Because integrin-linked kinase (ILK), a multidomain FA protein, is critically involved both in the adhesion of cells to the ECM by interacting with integrin β1 and in signal transduction in a PI3K-dependent manner, we hypothesized that ILK is involved in LN-2 signaling of oligodendrocyte cell spreading and myelination. To examine
whether ILK is expressed in oligodendrocytes, we double stained for ILK and NG2 as a marker for OPCs or MBP for mature oligodendrocytes (Fig. 7 A, arrows). Double staining clearly shows that ILK is expressed principally in interfascicular oligodendrocytes.

Next, we investigated ILK expression in vivo during early myelination of rat PNS SN (P days 1–5) and rat CNS ON (P7, P9–11, and P15) and in the CC and the ON of 5-wk control and dy/dy mice (Fig. 7 B). ILK was enriched in the rat PNS, specifically in cytoplasmic processes of Schwann cells at nodes of Ranvier. In the rat CNS ON, ILK expression was restricted to interfascicular oligodendrocytes (Fig. 7 B, arrows). Sections double labeled with ILK and Caspr, an axonal component of paranodal axoglial junctions, showed that ILK protein was detected only in oligodendrocytes from P9 onward, when myelination was underway. Double staining with ILK and Caspr in the CC and the ON of 5-wk control and dy/dy mice showed that ILK in the adult mouse CNS, as in the developing rat CNS, was localized to oligodendrocyte cell bodies (Fig. 7 C, arrows). There was no significant difference in the ILK expression pattern between control and dy/dy mice, whereas there appears to be a difference in Caspr immunoreactivity, suggesting paranodal abnormalities. Bars, 10 μm.

ILK shows colocalization with FA proteins such as integrin β1 and paxillin in oligodendrocytes

Given the reported role of ILK in adhesion complexes (Li et al., 1999), we labeled oligodendrocytes for ILK and the FA proteins integrin β1 and paxillin (Fig. 8 A). ILK was localized in the cell body and throughout the processes including their tips, with a high concentration of ILK at the growth cone–like termini. Digitally merged images of differentiating oligodendrocytes on LN-2 demonstrate that ILK colocalizes with integrin β1 in cell bodies and at FA sites in processes (Fig. 8 A, arrows).

Paxillin, a 68-kD FA adaptor molecule involved in integrin signaling, functions in regulation of the actin cytoskeleton (Turner, 2000). Like integrin β1, paxillin colocalized with ILK in FAs (Fig. 8 A). We evaluated the interaction between ILK and paxillin also by coimmunoprecipitation of ILK with paxillin from lysates of cells grown on BSA, LN-2, and in suspension (Fig. 8 B). The results suggest that the ILK–paxillin interaction is constitutive rather than adhesion induced.

LN-2 stimulates ILK activity and DN-ILK blocks LN-2-induced myelin membrane formation

To determine the functional role of ILK in LN-2–induced oligodendrocyte cell spreading, we used a dominant negative ILK
loss of function mutation in ILK (DN-ILK; kinase inactive), in which Glu 359 was mutated to Lys (Persad et al., 2000). When GFP-vector or GFP-DN-ILK was inserted into cells by adenovirus-mediated gene transfer, 60% of cells expressed GFP (unpublished data). Transfectants were replated on LN-2 in serum-free media and cultured for 1, 2, or 6 h. ILK activity was determined with exogenous MBP as a substrate. LN-2 induced ILK to phosphorylate MBP in GFP-transfected cells, with a peak at 1 h, but not in cells expressing DN-ILK (Fig. 9A). Furthermore, LN-2 stimulated ILK activity more effectively than any of the other ECMs at 1 h (Fig. 9B, P < 0.5).

ILK also can phosphorylate PKB/AKT on ser473 directly or indirectly (Delcommenne et al., 1998; Troussard et al., 2003). DN-ILK blocked phosphorylation of AKT ser473 (Fig. 9C). DN-ILK also inhibited phosphorylation of GSK-3β on ser 9, a downstream substrate of ILK (unpublished data). To test directly whether ILK is involved in LN-2–enhanced oligodendrocyte cell spreading, DN-ILK was expressed in OPCs, which were plated on LN-2 and incubated with vector or DN-ILK (10 or 20 MOI) for 3 d. As a control, we measured cell survival through the Live/Dead Cytotoxicity assay and found no significant cell death in DN-ILK (10 MOI) compared with vector (10 MOI), but, at the higher DN-ILK concentration of 20 MOI, we did find significant cell death (Fig. 9F).

Staining for O1 and DAPI revealed that 50% of GFP-transfected O1+ cells (10 or 20 MOI) formed broad sheetlike processes consistent with myelin membrane sheets (Fig. 9, D and E). In DN-ILK transfected cells, formation of these sheetlike processes was decreased by nearly 80% (P < 0.001). Only 10% of cells transfected with 10 MOI DN-ILK formed membrane sheets, and an even smaller percentage with 20 MOI DN-ILK (Fig. 9, D and E). These results demonstrate that ILK is necessary for LN-2 signaling in oligodendrocyte cell spreading and myelin formation. In summary, LN-2–induced oligodendrocyte cell spreading and myelination in CNS occurs in a PI3K/ILK-dependent manner.

Discussion
LN-2 is required for normal myelination in the CNS
Although congenital LN-2 deficiency in humans leads to CNS white matter changes as detected by magnetic resonance imaging as well as peripheral neuropathy and muscular dystrophy (Morandi et al., 1999), dy/dy mice with spontaneously occurring mutations in LN-2 were reported to exhibit phenotypic abnormalities limited to PNS myelination and skeletal muscle (Bradley and Jaros, 1979), except for the report of CNS myelin abnormalities in mice by Tsuji and Matsushita (1985). Here, we analyzed 5-wk-old dy/dy and littermate control mice for possible involvement of this gene in CNS myelination. In the CC of dy/dy mice, we found (1) reduced thickness of the CC; (2) a 30% reduction in the concentration of mature oligodendrocytes without a...
change in total glial cell concentration; and (3) defects in myelin lamellar compaction and sheath thickness. These results agree with previous studies (Buttery and ffrench-Constant, 1999) showing that LN-2 promotes formation of myelin membrane in vitro and further suggest that dy/dy mice may show developmental defects expressed as a disturbance in lamellar organization or oligodendrocyte cell volume. However, these results may also reflect a decrease in oligodendrocyte survival.

Small-sized axons are preferentially affected
Hypomyelination is particularly evident in small-sized axons. We found an $\sim 30\%$ reduction in the concentration of small-sized (<$0.5 \mu m$) axons that were myelinated in dy/dy CC, whereas the total number of myelinated axons in the same area was not changed. The preferential reduction in myelination of small-sized axons may reflect an inability of individual oligodendrocytes to extend multiple cytoplasmic processes and envelope a cluster of small-sized axons, without a corresponding defect in their ability to wrap around large-sized axons (Bjartmar et al., 1994). Oligodendrocyte cell bodies might lie closely associated with large-sized axons, without a requirement to reach them by extending cytoplasmic processes (Hildebrand et al., 1993). Accordingly, impaired formation of oligodendrocyte processes and membrane sheets would cause a more pronounced hypomyelination in tracts rich in small-sized axons than in tracts with large-sized axons. Another possible explanation for the differences between the abnormalities in small- and large-sized axons is that myelination may be delayed in the CNS of dy/dy mice, as it is in the PNS, because large-sized axons are generally myelinated earlier than small-sized axons in the CNS (Schwab and Schnell, 1989).

Regional differences
The dramatic differences between PNS and CNS myelination in the spontaneous dy/dy mutant mouse, described by several investigators approximately three decades ago (Bradley and Jaros, 1979), merit reconsideration. In segments of nerve roots of almost all cranial nerves and segments of both dorsal and ventral roots of cervical, lumbar, and sacral spinal nerves examined by EM, large clusters of small diameter axons are completely unmyelinated and lie contiguous with one another, without intervening Schwann cell cytoplasmic strands. Given that the gene product, LN-2, is a surface membrane constituent of axons, it now seems clear that the myelination problem at these restricted sites results not from a deficient number of Schwann cells caused by reduced cell division or excessive cell death, but from a failure of myelinating cells in the PNS and CNS to respond to the β1 component of LN-2 via PI3K and ILK by extending cytoplasmic membrane sheets is shown. 20 MOI DN-ILK decreases cell spreading up to 75% compared with vector alone ($P < 0.005$, asterisks).

(F) Cell survival was measured by the Live/Dead Cytotoxicity assay. There is no statistical difference in cell survival between 10 MOI vector and 10 MOI DN-ILK, whereas 20 MOI DN-ILK induces cell death ($P < 0.01$, asterisks). (E and F) Results are presented as ±SEM; and comparisons by ANOVA are significant (indicated by asterisks).
sheets, which are to envelop axons and form myelin lamellae. An unresolved problem is why myelination in the PNS is essentially normal in the few millimeters of the nerve roots closest to the SC, as well as throughout nerves distal to the roots, but grossly abnormal in relatively short intervening segments of the roots. This is a general problem with neurological genetic diseases, which commonly show a remarkable selectivity within what we generally assume to be a uniform population of target cells.

Here, we found regional differences in myelination and oligodendrocyte cell spreading within the CNS: (1) SC corticospinal tracts exhibited no reduction in myelin content and no dysmyelination, in contrast to the ON and CC, which were hypomyelinated and showed many dysmyelinated axons; and (2) Oligodendrocytes of the SC are less responsive to LN-2 than those of the CC. These data suggest that LN-2 might be a crucial factor for myelination only in selected areas of the CNS. Interestingly, Fyn knockout mice exhibit a pattern of regional differences within the CNS similar to what we have found in dy/dy mice (Sperber et al., 2001).

**LN-2 signaling in oligodendrocyte cell spreading is PI3K dependent**

We observed that LN-2, TSP-1, and TN-C all activated the phosphorylation of AKT at 45 min. However, for the anti-adhesive proteins, TSP-1 and TN-C, PI3K and AKT activation is transient. LN-2 activated AKT phosphorylation continuously for up to 6 h, suggesting that sustained AKT phosphorylation might be important for LN-2–induced cell spreading in oligodendrocytes. A role for AKT phosphorylation is also supported by our data that DN-AKT expression decreased LN-2–induced cell spreading. However, longer-term AKT activation remains to be examined.

PI3K, but not MAPK, participates in LN-2–enhanced myelin membrane sheet formation in oligodendrocytes in vitro. Previous findings had suggested that PI3K is involved in the regulation of actin cytoskeleton organization during integrin-mediated cell spreading (Seger et al., 2001). We show that PI3K inhibitors at concentrations (0.01 μM wortmannin and 0.5 μM LY294002) that do not affect cell viability prevent the formation of the LN-2–induced giant lamellae characteristic of oligodendrocytes, but do not change the number of O1° cells. One reason why PI3K inhibitors might not affect cell survival on LN-2 is because LN-2 may switch in survival signaling, from PI3K to MAPK pathway in mature oligodendrocytes (Colognato et al., 2002).

Our results are supported by previous data showing that TSP-1 influences the migration (Scott-Drew and French-Constant, 1997) and TN-C promotes proliferation (Garcion et al., 2001), but not differentiation through αvβ1 and αvβ3, respectively, in oligodendrocytes. In our short-term culture system, this LN-2–enhanced myelin membrane formation was not related to cell survival. However, previous studies have reported that LN-2 can function as a survival cue for mature oligodendrocytes (Colognato et al., 2002), and we cannot rule out the possibility that LN-2 in long-term culture might support oligodendrocyte cell survival.

**ILK plays a critical role in oligodendrocyte cell spreading**

Here, we show ILK expression in the CNS in vivo at high levels in interfascicular oligodendrocytes, with a time course that parallels myelination (Domercq et al., 1999). Because LN-2 is expressed on axons at the time of myelination (Powell et al., 1998), ILK may play an important role in LN-2–induced CNS myelination. Also, in the SN of the PNS, ILK was most prominent in Schwann cell cytoplasm, especially at and near nodes of Ranvier. Consistent with the in vivo data, we find that ILK is enriched in FAs in oligodendrocytes in vitro. Recently, ILK-deficient mice have been reported to die at the peri-implantation stage because of abnormal accumulation of actin at sites of integrin attachment at the basement membrane zone, and ILK-deficient fibroblasts show impaired cell spreading and delayed formation of FAs (Sakai et al., 2003). This implies a role for ILK in regulation of the actin cytoskeleton, which might be implicated in oligodendrocyte cell spreading. Furthermore, paxillin is important in regulation of the actin cytoskeleton in Schwann cells through integrin B1 (Taylor et al., 2003) and ILK colocalizes with paxillin in FAs (Nikolopoulos and Turner, 2001). We found that ILK is colocalized with integrin B1 and is associated with paxillin in FAs of differentiated oligodendrocytes on LN-2.

We also show that LN-2 stimulates ILK activity in oligodendrocytes and that the inhibition of ILK by DN-ILK blocks myelin membrane formation induced by LN-2. The interaction of the kinase domain of ILK with integrin B1, paxillin and PKB/AKT suggests that integrin B1, paxillin and PKB/AKT might be important regulators of the ILK-mediated signaling pathway for LN-2–induced oligodendrocyte cell spreading. Another possible mechanism through which ILK might influence oligodendrocyte cell spreading is suggested by the multidomain structure and FA localization of ILK, which imply that ILK might function as a scaffolding protein mediating multiple protein–protein interactions (Tu et al., 2001).

**A distinction between hypomyelination and dysmyelination**

Maturation of oligodendrocytes is a complex process that we have divided, in an attempt to unravel a temporal series of molecular control mechanisms, into arbitrary “stages”, which include cell spreading, axonal ensheathment, and myelin compaction. We have shown that the dy/dy mouse has a reduced number of mature oligodendroglial cells and features hypomyelination and dysmyelination in several CNS tracts. ILK appears to be involved in the LN-2/integrin signaling that promotes the early maturation events of cell spreading and process formation. Hypomyelination is an expected consequence, but the basis for the dysmyelination component of the dy/dy phenotype remains to be elucidated. Many additional genes are also involved in oligodendrocyte maturation. Among them, the nonreceptor-type tyrosine kinase Fyn, a member of the src family, has been identified as a signaling molecule downstream of L-MAG (Umehori et al., 1994). Analysis of Fyn−/−, MAG−/−, and Fyn/MAG double knockout mice indicates that Fyn is involved in oligodendrocyte membrane and process formation, and MAG
in the formation of compact CNS myelin (Biffiger et al., 2000). There may be direct or indirect interactions of the ILK–LN-2–integrin components and the related FA proteins with Fyn, MAG, and additional molecular mediators.

### Materials and methods

#### Animals

Homozygous 129Re/dy/dy mice and control littermates 5 wk old were purchased from The Jackson Laboratory. Dystrophic mice were distinguished from their littermates at 2–3 wk old, when they began to exhibit hindleg weakness. The +/+ and +/dy/littermates were pooled as controls.

#### Reagents

- LN-2, TN-C, and laminin-α2 chain (M-chain) antibody were obtained from CHEMICON International Inc.; PLO was obtained from Sigma-Aldrich; TSP-1 and ICN were obtained from Biomedicals; purified bSA was obtained from New England Biolabs, Inc.; Wortmannin and LY294002 (PI3 K inhibitors) were obtained from Calbiochem; mAbs against A2B5, O4, and O1 were obtained from American Type Culture Collection; monoclonal and polyclonal anti-ILK and monoclonal anti-integrin β1, antipaxillin, and anti-β-tubulin antibodies and MBP were obtained from Upstate Biotechnology; antibody against MBP SM999 was obtained from Stemberger Monoclonals; polyclonal antibodies against phospho-AKT ser473, AKT, phospho–EGFK-ser9, and UO126 (MAPK inhibitor) were obtained from Cell Signaling; monoclonal CC1 antibody was obtained from Oncogene Research Products; secondary antibodies were obtained from Jackson Immunoresearch Laboratories; Alamar blue assay kit was obtained from Trehk Diagnostics; and the Live/Dead viability/Cytotoxicity assay kit was obtained from Molecular Probes.

#### Primary oligodendrocyte cultures

Primary cultures of OPCs were prepared from the cerebral hemispheres of 2-d-old Sprague-Dawley rats as described previously (Vartanian et al., 1997). Purified OPCs were plated on PLO-coated 60-mm dishes and incubated in the presence of 10 ng/ml PDGF and basic FGF for 3 d. To produce differentiated oligodendrocytes, cells were plated on different ECM substrates after trypsinization for 8–10 min and incubated in serum-free media with N2 supplement (N2) for another 3 d. Various ECMs, 10 µg/ml bSA, 1 µg/ml LN-2, 15 µg/ml TSP-1, and 3 µg/ml TN-C were incubated in PLO-coated 24-well dishes at 37°C for 1 h and washed three times with PBS.

#### Immunofluorescence microscopy

Immunofluorescence staining for A2B5, O4, or O1 was performed as described previously (Park et al., 2001). For the staining of FA proteins such as ILK, integrin β1, or paxillin, cells plated on LN-2 were extracted with 0.5% Triton-X 100 buffer before fixation. ONs from rats and control and dy/dy mice were dissected and immediately fixed in ice cold 4% PFA in 0.1 M cacodylate buffer, pH 7.4, and incubated for 1.5 h in 1% osmium tetroxide/1.5% potassium ferrocyanide, washed in H2O, followed by maleate buffer, pH 5.2, and incubated for 1 h in 1% uranyl acetate in maleate buffer, pH 5.1. Blocks were washed in 0.1 M cacodylate buffer, pH 7.4, and incubated for 1 h in 1% uranyl acetate in maleate buffer, pH 5.1. Sections were dehydrated in graded ethanols, followed by propylene oxide, and embedded in TAAB Epon. Sections for light microscopy were cut at 0.5 µm and stained with alkaline toluidine blue. Ultrathin sections for EM were stained with uranyl acetate and lead citrate, and examined with a transmission electron microscope (model 1200EX; JEOL).

#### Immunohistochemical staining

Mice were perfused with 4% PFA and held overnight in the same fixative. Segments of brain were embedded in paraffin and sectioned at 6 µm. The total cell concentration was counted in sections stained with H&E. Mature oligodendrocytes in the CC were identified in 6 µm paraffin sections incubated at 4°C overnight with CC1 (1:3). CC1 immunoreactivity was detected with the ABC Elite Kit for mouse (Vector Laboratories) with DAB substrate.

#### EM

Mice were anesthetized with CO2 gas and perfused transcardially with 0.9% NaCl, followed by 2% PFA and 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.4. Brain, ON, and cervical SC were then removed, trimmed, and held overnight in the same fixative. Tissue blocks were washed in 0.1 M cacodylate buffer, pH 7.4, and incubated for 1.5 h in 1% osmium tetroxide/1.5% potassium ferrocyanide, washed in H2O, followed by maleate buffer, pH 5.2, and incubated for 1 h in 1% uranyl acetate in maleate buffer, pH 5.1. Blocks were washed in maleate buffer, dehydrated in graded ethanols, followed by propylene oxide, and embedded in TAAB Epon. Sections for light microscopy were cut at 0.5 µm and stained with alkaline toluidine blue. Ultrathin sections for EM were stained with uranyl acetate and lead citrate, and examined with a transmission electron microscope (model 1200EX; JEOL).

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