Glial and neuronal isoforms of Neurofascin have distinct roles in the assembly of nodes of Ranvier in the central nervous system

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Introduction

Nodes of Ranvier are the sites of action potential propagation in myelinated fibers, and their formation is essential for the switch to rapid nerve impulse transmission in the developing vertebrate nervous system. Axonal protein complexes containing voltage-gated sodium channels are assembled at nodes of Ranvier in response to myelination. In addition to sodium channels, components of the nodal complex include BiV-Spectrin, AnkyrinG, Contactin (in the central nervous system [CNS]), NrCAM (in the peripheral nervous system [PNS]), and a neuronal isoform of Neurofascin, Nfasc186 (Davis et al., 1996; Berghs et al., 2000; Jenkins and Bennett, 2002; Yang et al., 2004). The paranodal axoglial junctions that flank the node are formed by an adhesion complex between the glial isoform of Neurofascin, Nfasc155, and the axonal proteins Caspr (also known as Paranodin) and Contactin (Menegoz et al., 1997; Peles et al., 1997; Tait et al., 2000; Bhat et al., 2001; Boyle et al., 2001; Kazarinova-NOyes et al., 2001; Charles et al., 2002; Sherman and Brophy, 2005; Sherman et al., 2005). Axoglial junctions restrict the diffusion of nodal complexes but cannot cluster them in the PNS in the absence of Nfasc186 in vivo (Rios et al., 2003; Sherman et al., 2005).

Studies on myelinating cocultures have indicated a key role for nodal Nfasc186 in the clustering and stabilization of macromolecular assemblies at the CNS node of Ranvier (Lambert et al., 1997; Bennett and Lambert, 1999; Lustig et al., 2001; Koticha et al., 2006; Dzhashiashvili et al., 2007), and studies in vivo have supported this view (Sherman et al., 2005). Gliomedin, a protein expressed in the region of Schwann cell microvilli, may anchor Nfasc186 at the CNS node by virtue of its ability to interact with both Neurofascin and NrCAM (Eshed et al., 2005; Eshed et al., 2007; Maertens et al., 2007). However, there are indications that the mechanisms of node assembly may be different in the CNS. Soluble factors secreted by oligodendrocytes have been shown to promote sodium channel clustering in CNS axons in culture in the absence of axoglial contact (Kaplan et al., 1997). More recent work has refined this picture by showing that Na1,2α subunits cluster under the influence of a secreted factor; however, clustering of Na1,6α subunits, which are more characteristic of mature nodes, requires ensheathment by oligodendrocytes (Boiko et al., 2001;
Kaplan et al., 2001). These studies have further suggested that vesicular traffic is required to deliver sodium channels to nascent nodes (Kaplan et al., 2001).

Analysis of mice with mutations that affect the integrity of paranodal axoglial junctions has suggested that these structures, though not essential for the initial clustering of sodium channels, may be important for the long-term maintenance of the nodal complex (Bhat et al., 2001; Ishibashi et al., 2002; Rasband et al., 2003; Rios et al., 2003; Dupree et al., 2005). Derangement of the node after loss of axon–glial contact in tissue from multiple sclerosis patients caused by inflammatory damage to CNS axons is consistent with a role for the junction in stabilizing the node (Howell et al., 2006). However, thus far it has not been possible to discriminate between the relative contributions of nodal and paranodal Neurofascins in the initial assembly of the nodal complex in the CNS.

In this paper, we have studied how axonal and glial Neurofascins contribute to the clustering of sodium channels at CNS nodes of Ranvier in vivo by using Nfasc−/− mice with disrupted nodal and paranodal protein complexes. By selectively expressing either Nfasc186 or a truncated version of Nfasc155 by transgenesis in Neurofascin-null mice, we have been able to show that, in marked contrast to the PNS, each of the Neurofascin isoforms can independently rescue the CNS node of Ranvier. Furthermore, we have revealed an important function for the paranodal axoglial junction in promoting the migration and convergence of myelinating processes along axons. We propose that these two Neurofascin isoforms cooperate to assemble and maintain functional nodes of Ranvier in the vertebrate CNS.

Results

CNS phenotype of Nfasc−/− mice

Neurofascin-null animals die suddenly at 7 d after birth (Sherman et al., 2005). Therefore, analysis of their phenotype was performed at postnatal-day (P) 6. For most of this work, we used either teased fiber preparations or longitudinal sections from the ventral funiculus of the cervical spinal cord because we found that CNS myelination was most advanced there at this early stage of postnatal development in the mouse. Furthermore, this region of the spinal cord has relatively large axons, thus facilitating the preparation of teased fibers for microscopy and quantitation (Arroyo et al., 2002). Immunofluorescence showed that in Neurofascin-null nerves, there was extensive disruption to both the nodal complex (sodium channels, βIV-Spectrin, and AnkyrinG) and the paranodal axoglial junction (Caspr; Fig. 1 A).

The paranodal myelin marker Claudin 11 was used to identify paranodes (and by inference nodes; Gow et al., 1999). Contactin was present at both the nodes and paranodes of wild-type nerves (Fig. 1 A, figure and inset), as previously reported (Kazarinova-Noyes et al., 2001) but was no longer detectable at either site in the mutant (Fig. 1 A). Similarly, AnkyrinG is not only a major component of the node of Ranvier, but it is also detectable at wild-type paranodes at this stage of CNS development (Rasband et al., 1999; Jenkins and Bennett, 2002); however, it is lost in the mutant in both locations (Fig. 1 A).

Disruption of the junctional complex at the paranode in the CNS was accompanied by loss of the characteristic septate junctions between the paranodal loops and the axolemma (Fig. 1 B). However, although it was mislocalized, there were normal amounts of Caspr in mutant spinal cord tissue, suggesting that disruption of the axoglial adhesion complex did not impair the biosynthesis and/or stability of the protein (Fig. 1 C).

Electron microscopy revealed myelinating and unmyelinated profiles in both wild-type and mutant nerves (Fig. 2 A). However, Western blotting showed that the levels of the myelin proteins myelin-associated glycoprotein (MAG) and myelin basic protein (MBP) in the CNS of the mutant at P6 were reduced compared with wild-type tissue and that they were similar to those observed at P4 in the wild-type (Fig. 2 B). This suggested...
that myelination might be either retarded or halted in the mutant. The reduction in myelin protein in the mutant was accompanied by a significant decrease in the number of myelinated axons observed at P6 in the ventral funiculus of the mutant cervical spinal cord (Fig. 2 C). The reduction in the amount of myelin was not a consequence of a general decrease in the number of axons in the CNS of Neurofascin-null animals because the total number of axons in the optic nerves of mutants was normal (Fig. 2 D). Neither was the reduced extent of myelination a consequence of an inadequate pool of myelin-forming glia because the numbers of oligodendrocytes in wild-type and mutant cervical spinal cord sections were not significantly different (Fig. 2 E).

**Nfasc186 rescues the nodal complex**

We used a transgenic approach to ask if expression of Nfasc186 FLAG tagged at its C terminus could reconstitute the nodal complex in the CNS axons of Nfasc−/− mice in vivo in the absence of an intact axoglial adhesion complex. First, it was important to demonstrate that this epitope-tagged protein was targeted appropriately to the node of Ranvier and that it colocalized with the nodal complex (sodium channels, BIV-Spectrin, Contactin, and AnkyrinG) in Nfasc−/− mice (Fig. 3 A). Then, we showed that expression of transgenic Nfasc186 on a Neurofascin-null background was fully able to rescue the nodal complex (Fig. 3 A). Intriguingly, and in contrast to Contactin, AnkyrinG in the rescued fibers resumed its normal localization at both the node and paranode (Fig. 3 A). This suggests that the transient expression of AnkyrinG at the paranodes during the postnatal development of myelinated CNS fibers does not depend on either Nfasc155 or its axonal partners Caspr and Contactin because in the absence of Nfasc155, there was no amelioration of the mislocalization of Caspr (Fig. 3 B). The transient nature of paranodal AnkyrinG is shown by its loss in both wild-type and Nfasc186-rescued nerves by P16 (Fig. 3 D).

Importantly, the nodal rescue by Nfasc186 was functionally significant because these mice survived beyond P7 only to succumb at P18–19. This allowed us to study two developmentally regulated events in myelinated CNS axons, namely, the expression of juxtaparanodal potassium channels and the upregulated switch to sodium channels of the Nav1.6 type. Absence of an intact axoglial junction caused juxtaparanodal potassium channels of the Shaker type Kv1.1 to redistribute to ectopic sites close to the nodal sodium channels, as has been observed in other mutants with deranged axoglial junctions (Fig. 3 B; Bhat et al., 2001; Boyle et al., 2001). However, the developmentally regulated switch to sodium channels of the Nav1.6 type was unimpaired by the absence of an axoglial junction (Fig. 3 C) as was found in the Jimpy mouse (Jenkins and Bennett, 2002).

We have shown previously that a form of Nfasc155 with most of the C terminus replaced by a FLAG tag (Nfasc155ΔIC) is targeted to the paranodal loops of myelinating Schwann cells and oligodendrocytes (Sherman et al., 2005). Further, we showed that it can reconstitute the axoglial junctional complex in the peripheral nerves of Neurofascin-null mice (Sherman et al., 2005), presumably by virtue of the ability of the extracellular domain to interact with components of the axonal Caspr–Contactin complex (Charles et al., 2002; Gollan et al., 2003). However, these reconstituted paranodal complexes could not rescue sodium channels in the absence of Nfasc186 at PNS nodes (Sherman et al., 2005). Using the neuron-specific neurofilament light chain (NF-L) promoter, Nfasc186 transgenic mice also displayed robust expression in the PNS, which allowed us to show directly in vivo that Nfasc186 could rescue the nodal complex (sodium channels, BIV-Spectrin, AnkyrinG, and NrCAM) in the absence of an intact axoglial junction in PNS axons (Fig. 4 A). Interestingly, we were also able to show that NrCAM is a component of PNS nodes but that it is also absent from CNS nodes (Fig. 4 B).

**Reconstitution of the axoglial adhesion complex rescues CNS nodal sodium channels**

Although reconstitution of the axoglial junction cannot rescue the nodal complex in the PNS in the absence of Nfasc186, we wished to ask if the same holds true in the CNS. Nfasc155ΔIC
reconstituted the paranodal axoglial adhesion complex in the CNS of Nfasc<sup>−/−</sup> mice (Fig. 5 A), but, in complete contrast to the PNS, the correct localization of nodal sodium channels was also rescued (Fig. 5 B). Other components of the CNS nodal complex, such as βIV-Spectrin, were also correctly localized (Fig. 5 B). Unlike the Nfasc186 rescue, mice expressing Nfasc155ΔIC on a Neurofascin-null background did not have an enhanced life expectancy. This may be because of the fact that nodal complexes in the PNS are not rescued concomitantly with those in the CNS.

Nodal gaps of less than 5 μm in width were identified by symmetrical Claudin 11 immunostaining, and the percentage of these that were immunostained for sodium channels was measured. In wild-type axons of the spinal cord ventral funiculus, 100% of such nodal gaps immunostained for sodium channels (Fig. 5 C). The efficiency of sodium channel rescue was equally high when reconstituting the axoglial junction with Nfasc155ΔIC or when expressing Nfasc186 (91 ± 4 and 96 ± 2%, respectively; mean ± SEM; n = 3 each; Fig. 5 C). In both cases, there was no significant difference in the frequency of sodium channel staining when compared with wild-type nodes (Fig. 5 C).

Interestingly, it was possible to detect sodium channels at a minority of Claudin 11–flanked nodes in Neurofascin-null nerves (16.0 ± 6.1%; mean ± SEM; n = 3; Fig. 5 C). This suggested that sodium channels could still be targeted to the node in Nfasc<sup>−/−</sup> mice but that the efficiency of their delivery and/or their stability upon arrival was impaired in the absence of Nfasc186 or an intact axoglial junction. The ability of the Nfasc155–Caspr–Contactin adhesion complex to promote the concentration of voltage-gated sodium channels at CNS nodes in the absence of Nfasc186 is in marked contrast to the PNS node (Sherman et al., 2005), and this difference was confirmed in this paper using a transgene encoding the complete Nfasc155 protein (Fig. 5 D).
that it was able to form Nav1.6 channel clusters at nodes but that these became more diffuse and less concentrated with time (Rasband et al., 2003). Interestingly, Contactin and Caspr were still detectable in paranodal complexes in these mice, whereas Nfasc155 was not (Schafer et al., 2004). These observations may be reconciled by our demonstration here that either Nfasc186 or Nfasc155 (through the formation of the axoglial complex) can promote the assembly of the nodal complex. Thus, in the case of the ceramide galactosyltransferase mutant mice, Nfasc186 delivered to the axolemma, possibly guided by local concentrations of Caspr and Contactin, might drive the assembly of the node in the absence of the paranodal junction.

The tips of oligodendrocyte processes visualized in the teased fiber preparations used in the present work are enriched in Nfasc155 and Caspr. The early formation of axoglial junctions has been observed previously in the optic nerve, and it is known that axoglial septate junctions can be detected in the same nerve as early as after one turn of the myelinating process around an axon (Wiggins et al., 1988; Rasband et al., 1999). The fact that process migration is retarded in the absence of these proteins suggests that the axoglial adhesion complex has a role in promoting their extension. Nevertheless, the ability of the processes to ultimately converge shows that the axoglial complex is not solely responsible for allowing oligodendrocyte processes to extend. Recent evidence for the involvement of the WAVE proteins and their associated links with the actin cytoskeleton in oligodendrocyte process extension provides a possible additional mechanism (Kim et al., 2006).

Signaling via the axoglial adhesion complex might stimulate oligodendrocytes to extend their myelinating processes. However, Nfasc155 lacking most of its C terminus can not only reconstitute the adhesion complex but can also cluster nodal proteins, which shows that intracellular signaling mediated via the carboxy terminus of Nfasc155 is not necessary to promote the assembly of the node of Ranvier in the CNS. This raises the intriguing idea that intracellular signaling in the axon in response to axoglial interaction, possibly mediated by the cytoplasmic

Discussion

The mechanisms of assembly of the node of Ranvier and the clustering of sodium channels are likely to occur by different mechanisms in the CNS when compared with the PNS, not least because of the different structure and protein composition of their nodal environments (Kaplan et al., 1997, 2001; Poliak and Peles, 2003; Salzer, 2003; Sherman and Brophy, 2005). For example, Contactin is a characteristic component of CNS nodes but is absent from PNS nodes, whereas we find that the reverse is true for NrCAM (Kazarinova-Noyes et al., 2001).

Previous work has provided strong circumstantial evidence that the axoglial junction might have a role in the formation of the CNS node. Analysis of the developing optic nerve showed that Caspr staining, which is indicative of the formation of axoglial complexes, preceded the detection of sodium channel clusters, which were initially typically broad (Rasband et al., 1999). The same authors showed that Shiverer mutant mice lacking normal axoglial junctions had reduced and ectopic sodium channel clusters. This indicated, probably for the first time, that there was a relationship between the integrity of the axoglial junctional complex and the ability of myelinated CNS axons to cluster sodium channels. Subsequent analysis of another mutant, UDP-galactose/ceramide galactosyltransferase–deficient mice, which also lacks intact paranodal axoglial junctions, showed that it was able to form Nav1.6 channel clusters at nodes but that these became more diffuse and less concentrated with time (Rasband et al., 2003). Interestingly, Contactin and Caspr were still detectable in paranodal complexes in these mice, whereas Nfasc155 was not (Schafer et al., 2004). These observations may be reconciled by our demonstration here that either Nfasc186 or Nfasc155 (through the formation of the axoglial complex) can promote the assembly of the nodal complex. Thus, in the case of the ceramide galactosyltransferase mutant mice, Nfasc186 delivered to the axolemma, possibly guided by local concentrations of Caspr and Contactin, might drive the assembly of the node in the absence of the paranodal junction.

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Figure 4. Nfasc186 rescues the PNS nodal complex in Nfasc<sup>−/−</sup> mice. [A] FLAG-tagged Nfasc186 can rescue the nodal complex in teased sciatic nerve fibers immunostained for the nodal proteins voltage-gated sodium channels (Nav), βIV-Spectrin, AnkyrinG, and NrCAM. Myelin was visualized using MBP as a marker. [B] NrCAM is present at PNS nodes but absent from CNS nodes. Bars, 5 μm.

Neurofascin-null mice rescued with the Nfasc186 transgene had an extended lifespan, and this allowed us to ask whether oligodendrocyte process migration was either delayed or was completely stalled in the absence of Nfasc155 at P6. In contrast to P6 (Fig. 6 F), at P16 there was no significant difference in the interheminodal gap between Nfasc<sup>−/−</sup> and Nfasc<sup>−/−</sup>/Nfasc186 mice (Fig. 6 G). Hence, the absence of an axoglial adhesion complex delayed, rather than prevented, the extension and convergence of oligodendrocyte processes altogether, and electron microscopy showed that this delay did not cause dysmyelination (Fig. 6 H).
Figure 5. Reformation of the paranodal adhesion complex rescues the node. (A) Immunostaining of longitudinal sections of the spinal cord ventral funiculus in the CNS shows that FLAG-tagged Nfasc155ΔIC is targeted to the paranodes of myelinated axons and that it can rescue Caspr to the paranodal axoglial junctional complex in Nfas−/− mice. (B) Rescue of Caspr to the paranode by Nfasc155ΔIC on a Neurofascin-null background reconstitutes voltage-gated sodium channels (Nav) and βIV-Spectrin to the nodal complex. (C) In the CNS at P6, both Nfasc186 and Nfasc155ΔIC are equally effective at reconstituting nodes with sodium channels (Nav) when expressed in the absence of endogenous Neurofascins. The percentage of nodal gaps of <5 μm in width identified by Claudin 11 immunostaining that also had sodium channels was measured. (D) Full-length Nfasc155 tagged with FLAG at the C terminus reconstitutes the axoglial junction in both CNS and PNS, but nodal Nav channels are only rescued in the CNS. Neurofascins were detected in the wild type using a pan anti-Neurofascin antibody that recognizes both Nfasc155 and Nfasc186. Values are means ± SEM. n = 3 mice for each condition. Bars, 5 μm.

The internodes of CNS axons are up to three times shorter than those in the PNS at a given axon diameter. Nevertheless, the CNS node of Ranvier still accounts for <1% of the length of the adjacent internode (Ramon y Cajal, 1909; Fraher, 1978). In spite of the fact that nodes constitute such short segments of the axon, clustered sodium channels are still detectable at ~16% of nascent nodes in Nfas−/− nerves. This indicates that sodium channels are not randomly delivered to the axolemma but are targeted to the nascent node independently of either Nfasc186 or Nfasc155, although one or both of these proteins is clearly required to stabilize the nascent node. Interestingly, a previous study found that ~12% of sodium channel clusters in the developing optic nerve was not associated with Caspr-positive axoglial junctions (Rasband et al., 1999). However, these clusters would be expected to contain Nfasc186, which, as the present work shows, would be sufficient to promote the assembly of the nodal complex.

Sodium channels are probably delivered to the nascent node via an exocytic pathway (Kaplan et al., 2001). It has been proposed that AnkyrinG might regulate such targeting (Jenkins and Bennett, 2002); however, this still leaves the question open as to how AnkyrinG is itself delivered to the nascent node. Once the sodium channels with their associated β subunits reach the nodal membrane, they would be expected to interact with Nfasc186 and βIV-Spectrin to form a macromolecular complex (Ratliffe et al., 2001; McEwen and Isom, 2004). In the absence of Nfasc186, the complex would diffuse in the lateral plane of the membrane, and without an intact axoglial junction to function as a sieve or barrier to their further dispersal, the nodal proteins may be endocytosed or proteolyzed in the internodal region, as occurs in the PNS (Pedraza et al., 2001; Dzhavashvili et al., 2007). Reintroduction of Nfasc186 would reduce the dissipation of the complex by anchoring it at the node. Because sodium channels, their β subunits, and AnkyrinG can interact with Nfasc186 directly, their presence at the node could also drive the targeted delivery of Nfasc186 (Ratliffe et al., 2001; Jenkins and Bennett, 2002; McEwen and Isom, 2004).
The clustering of sodium channels can occur independently of the presence of an intact axoglial junction. Nevertheless, there is a tendency for nodal sodium channels to disperse over time when the paranodal junctional complex is disrupted (Rios et al., 2003). The importance of an intact axoglial junction for the long-term stability of the nodal complex has also been demonstrated in mouse mutants and after chemical disruption of the myelin sheath (Ishibashi et al., 2002; Dupree et al., 2005). The limited extension of viability in Neurofascin-null mice expressing transgenic Nfascl86 may reflect their lack of intact axoglial junctions because Caspr-null mice lacking paranodal junctions in the CNS and PNS generally die between P21 and P33 (Bhat et al., 2001). Interestingly, the tendency for the nodal complex to become more diffuse is more pronounced in the CNS compared with the PNS (Rios et al., 2003). This suggests that the junctional complex plays a more important role in containing the nodal complex in the CNS than in the PNS, which is consistent with a more prominent role for the axoglial junction in assembling the CNS nodal complex.

An intriguing question is why CNS nodes exploit two distinct mechanisms to concentrate the macromolecular complex required for saltatory conduction at the node of Ranvier, whereas in the PNS, Nfascl86 has a uniquely important role. The role of Nfascl86 in the PNS had been inferred from previous experiments both in vivo and in vitro (Sherman et al., 2005; Dzhashiashvili et al., 2007), and here we have demonstrated its absolute requirement in PNS node assembly in vivo. Once anchored at the PNS node via extracellular interactions, possibly mediated by gliomedin and intracellular association with the actin cytoskeleton by means of AnkyrinG and βIV-Spectrin, Nfascl86 participates in an exceptionally stable complex, as indicated by the relative insensitivity of sodium channels to dispersion after disruption of the paranodal axoglial junction (Bhat et al., 2001; Boyle et al., 2001; Rios et al., 2003). The uniquely PNS nodal component, NrCAM, may also play a part here because it is known to interact with Nfascl86 (Volkmer et al., 1996). In the absence of NrCAM, CNS nodes may depend on the axoglial junction to provide an independent mechanism for focusing and concentrating sodium channels to complement their cis interactions with Nfascl86. Future application of the transgenic rescue strategy described in this work should allow us to functionally dissect the protein–protein interactions by which Nfascl86 and Nfascl55 cooperate in the assembly of the CNS node of Ranvier.

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Materials and methods

Animals

All animal work conformed to UK legislation (Scientific Procedures Act 1986) and to the University of Edinburgh Ethical Review Committee policy. The generation of Nfas*−/− and transgenic mice, Nfas IC, expressing a truncated form of Nfas 1.55 under the control of the Pip promoter has been previously described (Sherman et al., 2005). Transgenic mice, Nfas IC, expressing the full-length Nfas 1.55 with a FLAG tag at its C terminus under the control of the Pip promoter, were generated as for the Nfas IC transgenic line. Transgenic mice expressing a full-length DNA encoding Nfas 1.16 under the control of the NF-L promoter Nfas 1.16 were similarly generated by pronuclear injection. After introducing a FLAG tag sequence at the 3′ of the coding region, the DNA was cloned into the Clal site in the pCAG-L vector (J.P. Julien, Laval University, Quebec City, Canada) and D.E. Merry (Thomas Jefferson University, Philadelphia, PA) and was released using Ascl and Notl. After backcrossing to a C57BL/6 background, one of the lines was interbred with Nfas*−/− mice to generate Nfas −/− Nfas 1.16 mice.

Antibodies, microscopy, and Western blots

Unless indicated otherwise, all microscopic images were from nerves of 6-d-old animals. For immunostaining of CNS or PNS teased fiber preparations at room temperature, the ventral funiculus of the cervical spinal cord, 10 random images from three animals were captured at a 65K and relating this to the total area of the optic nerve using Openlab software. Electron microscopic images were captured using a Biowin (Philips) either by conventional photography or by using a digital camera (Orius; Gatan). Photographic negatives were scanned and digitized. All figures were prepared using Photoshop version 7.0 (Adobe) and were not subjected to any subsequent image processing. Western blotting was performed as previously described (Sherman et al., 2005).

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