Characterization of the Shark Myelin Po Protein

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Introduction

Myelin, in its compact form, is the insulating sheath that covers axons in the central and peripheral nervous system, allowing rapid nerve conduction. It consists of glial plasma membrane tightly wrapped around axons and devoid of any cytoplasmic fluid. Myelin compaction is dependent on the presence of highly adhesive molecules that keep the two sides of the membrane in tight contact. The Po glycoprotein (Po) is the major component of the peripheral nervous system (PNS) myelin of mammals. This protein has been shown to bind in a homophilic manner to an opposing membrane and is the molecule responsible for myelin compaction [D’Urso et al., 1990; Filbin et al., 1990; Giese et al., 1992]. The exact role that Po protein has played in the evolution of myelin is still unclear, but several phylogenetic observations suggest that it is a crucial component in the development of myelin as a multi-lamellar membrane structure. Sharks, which appeared in the fossil record about 400 million years ago, are the first fully myelinated organisms. In this study we investigated the expression pattern of shark myelin Po to suggest a way it might have played a role in the evolution of myelin in the central nervous system. We found that sharks have more than two isoforms (32, 28 and 25 kD), and that some of these might not be fully functional because they lack the domains known for Po homophilic adhesion.

Key Words
Development · Evolution nervous systems · Myelin · Shark · Developmental evolution · Elasmobranch · Embryology · Po protein

Abstract
Myelin, the insulating sheath made by extensive plasma membrane wrapping, is dependent on the presence of highly adhesive molecules that keep the two sides of the membrane in tight contact. The Po glycoprotein (Po) is the major component of the peripheral nervous system (PNS) myelin of mammals. The exact role that Po protein has played in the evolution of myelin is still unclear, but several phylogenetic observations suggest that it is a crucial component in the development of myelin as a multi-lamellar membrane structure. Sharks, which appeared in the fossil record about 400 million years ago, are the first fully myelinated organisms. In this study we investigated the expression pattern of shark myelin Po to suggest a way it might have played a role in the evolution of myelin in the central nervous system. We found that sharks have more than two isoforms (32, 28 and 25 kD), and that some of these might not be fully functional because they lack the domains known for Po homophilic adhesion.
branchs and teleost fish, Po is the major myelin protein component not only in the PNS but also in the central nervous system (CNS) [Waehneldt et al., 1986; Saavedra et al., 1989; Stratmann and Jeserich, 1995]. These observations have suggested to researchers that the transition between non-compact myelin to compact myelin parallels the appearance of Po in evolution.

Cloning shark Po revealed that this glycoprotein is conserved (~46%) throughout vertebrate evolution (fig. 1) and is the product of a single mRNA transcript [Waehneldt et al., 1987; Saavedra et al., 1989; Stratmann and Jeserich, 1995]. Furthermore, the cloning of a Po-like glycoprotein from trout CNS shows that it has about 50% sequence homology with shark and rat Po and also results
from one mRNA transcript [Stratmann and Jeseric, 1995]. Indeed, Po in elasmobranchs also carries the same HNK-1 carbohydrate epitope as Po in mammals [Zand et al., 1991].

In mammals only one Po isoform has been detected [Uyemura and Kitamura, 1991], whereas in sharks [Tai and Smith, 1983; Nunn et al., 1987; Saavedra et al., 1989] and chickens [Nunn et al., 1987] at least two isoforms are present. Bony fishes also seem to have two Po-like proteins (IP1 and IP2) [Stratmann and Jeserich, 1995]. But the true identity of these Po-like proteins in shark has yet to be defined by methods more sophisticated than merely the determinations of molecular weight and type of glycosylation.

The value of studying these well known myelin proteins in sharks derives from the position that cartilaginous fish hold in the evolution of myelin. Sharks, which appeared in the fossil record about 400 million years ago, are the first fully myelinated organisms [Bakay and Lee, 1966; Waehneldt, 1990]. Before them, other organisms, such as lampreys and earthworms, have glial membrane loosely wrapped around axons, but not true compact myelin [Bullock et al., 1984; Waehneldt et al., 1987]. Cartilaginous fish are considered to be the first ones to have a compact myelin [Kitagawa et al., 1993]. All this implies that in the evolution of myelin several factors converged: the presence of glial cells, presence of adhesive molecules and evolutionary advantage over non-compact myelin. Therefore, studying the spatio-temporal expression of the major myelin protein in shark will help in our understanding of the evolution of myelin. In this study we set out to investigate the expression pattern of shark myelin Po as a way of understanding its contribution to the evolution of myelin in the central nervous system.

**Materials and Methods**

**Po Antibodies**

Rabbit antibodies against Po were raised by injection of purified bovine Po with Freund's adjuvant (RabA antibody) or of Po peptides with Titer-Max (peptide antibodies). The polyclonal antibody PoCy2, that recognizes the cytoplasmic domain of mouse Po, has been described elsewhere [Gould et al., 1995]; it detects the full length Po protein. For the anti-Po peptide antibodies, two sequences from shark Po [Saavedra et al., 1989] and one from rat Po protein [Lemke et al., 1988] were selected to raise peptide antibodies (fig. 1). In order to raise antibodies that will be specific for shark only, we chose sequences where the similarity between shark and mouse was the lowest. Thus Western blots using PoEx or PoCy1 on rat myelin did not show any cross-reactivity, confirming that these new antibodies are specific for shark Po.

**Shark Nervous System Preparation**

Shark nervous system regions from adult spiny dogfish (*Squalus acanthias* L. 1758) from Woods Hole Marine Biological Labs (MBL) were rapidly dissected and frozen on dry ice. Following the procedure reviewed and approved by the Animal Welfare Committee at MBL, the animals were anesthetized in 1:10,000 parts of MS222 in sea water until they were insensitive to touch and were then killed by decapitation. The following regions were dissected: telencephalon (olfactory bulbs and cerebral hemispheres), diencephalon (epiphysis), mesencephalon (optic tectum), metencephalon (cerebellum), myelencephalon (medulla), and trigeminal nerves [Wischnitzer, 1993; Butler and Hodos, 2005]. Shark embryos collected from pregnant *Squalus acanthias* were staged based on fetal length as described in Ballard for shark embryos that are fully developed, although not yet fully grown [Ballard, 1993]. We collected and immediately froze brains from 4.0, 4.4, 9, and 22 cm embryos. Crude membrane fractions were prepared by homogenizing each region in 4 volumes of 1X SDS Laemmli sample buffer (2% SDS, 10% Glycerol, 62.5 mM Tris pH 6.8) in the presence of protease inhibitors (1 µg/ml leupeptin, 2 µg/ml antipain, 10 µg/ml benzamidine, 1 µg/ml pepstatin, 1 µg/ml aprotinin, and 100 µM PMSF). The protein content in each homogenate was estimated with a protein assay kit (Bio-Rad). Aliquots of the homogenates (depending on the figure, the amounts ranged between 2 and 10 µg of total protein in a 50 µl volume) were loaded onto 12% polyacrylamide minigels (which have a 4% stacking gel run at 70 mA and resolving gel at 35 mA), transferred to polyvinylidene fluoride (PVDF) membranes (at 0.6 A for 18 h at 4 °C) in the presence of 2% SDS, 10% Glycerol, 62.5 mM Tris pH 6.8 following transfer buffer conditions: 50 mM Na₂HPO₄ (anhydrous), 2 mM EDTA, 0.025% SDS that was buffered to pH 5.5 with concentrated HCl. Blots were immunostained with 1:3,000 dilutions of each different antibody. Shark Po was detected by secondary anti-rabbit antibody coupled either to alkaline phosphatase or horseradish peroxidase (1:20,000), and visualized by BCIP/NBT following manufacturer instructions (KPL), respectively.

**Myelin Preparation**

Myelin was isolated from the brains of spiny dogfish (*S. acanthias*) by conventional procedure [Norton and Poduslo, 1973]. The spinal cord was homogenized in 0.25 M sucrose with protease inhibitors (1 µg/ml leupeptin, 2 µg/ml antipain, 10 µg/ml benzamidine, 1 µg/ml pepstatin, 1 µg/ml aprotinin, and 100 µM PMSF) and the concentration was brought up to 1.4 M sucrose by adding 8 M urea, 10 µl/H₉₂O₂, and 100 µl/MBL were rapidly dissected and frozen on dry ice. Following the procedure reviewed and approved by the Animal Welfare Committee at MBL, the animals were anesthetized in 1:10,000 parts of MS222 in sea water until they were insensitive to touch and were then killed by decapitation. The following regions were dissected: telencephalon (olfactory bulbs and cerebral hemispheres), diencephalon (epiphysis), mesencephalon (optic tectum), metencephalon (cerebellum), myelencephalon (medulla), and trigeminal nerves [Wischnitzer, 1993; Butler and Hodos, 2005]. Shark embryos collected from pregnant *Squalus acanthias* were staged based on fetal length as described in Ballard for shark embryos that are fully developed, although not yet fully grown [Ballard, 1993]. We collected and immediately froze brains from 4.0, 4.4, 9, and 22 cm embryos. Crude membrane fractions were prepared by homogenizing each region in 4 volumes of 1X SDS Laemmli sample buffer (2% SDS, 10% Glycerol, 62.5 mM Tris pH 6.8) in the presence of protease inhibitors (1 µg/ml leupeptin, 2 µg/ml antipain, 10 µg/ml benzamidine, 1 µg/ml pepstatin, 1 µg/ml aprotinin, and 100 µM PMSF). The protein content in each homogenate was estimated with a protein assay kit (Bio-Rad). Aliquots of the homogenates (depending on the figure, the amounts ranged between 2 and 10 µg of total protein in a 50 µl volume) were loaded onto 12% polyacrylamide minigels (which have a 4% stacking gel run at 70 mA and resolving gel at 35 mA), transferred to polyvinylidene fluoride (PVDF) membranes (at 0.6 A for 18 h at 4 °C) in the following transfer buffer conditions: 50 mM Na₂HPO₄ (anhydrous), 2 mM EDTA, 0.025% SDS that was buffered to pH 5.5 with concentrated HCl. Blots were immunostained with 1:3,000 dilutions of each different antibody. Shark Po was detected by secondary anti-rabbit antibody coupled either to alkaline phosphatase or horseradish peroxidase (1:20,000), and visualized by BCIP/NBT following manufacturer instructions (KPL), respectively.

**Endo F Treatment**

An 80 µg sample of the shark (*S. acanthias*) spinal cord myelin fraction was boiled in 400 µl of EndoF lysis buffer (50 mM Tris pH 8.6, 25 mM EDTA, 0.1% SDS and 0.5% NP-40) in the presence of 2 µl of β-mercaptoethanol. A cocktail of protease inhibitors (1 µg/ml leupeptin, 2 µg/ml antipain, 10 µg/ml benzamidine, 1 µg/ml pepstatin, 1 µg/ml aprotinin, and 100 µM PMSF) was added once the mixture cooled to room temperature, and 1 µl...
(1 U/ml) of EndoF (Boehringer Mannheim) was added. The mixture was incubated overnight at 37 °C; 8 volumes of acetone were added to precipitate protein, and the mixture was cooled for 2 h at –20 °C. The proteins were pelleted by centrifugation at 5,000 rpm for 5 min in a Sorvall bench-top unit. Acetone was removed and once the precipitate was dry, the proteins were re-suspended in 30 µl of SDS sample buffer and 5 µg of total protein was run on the gel.

Immunohistochemistry

Chiloscyllium punctatum (Müller and Henle 1838) shark cases were harvested from the Long Beach Aquarium of the Pacific and the embryos collected at different developmental stages [Ballard, 1993]. The youngest collected embryos were 3 cm when they have developed most of their nervous system and they look like very small sharks; the oldest embryos were 10 cm, a stage at which they show a lot of physical activity and look like their adult counterparts. Embryos were fixed in Carnoy’s (70% ethanol, 20% formaldehyde and 10% glacial acetic acid) overnight at 4 °C and kept in 70% ethanol at –20 °C until histological preparation. In order to obtain good morphological results, the embryos needed very extensive dehydration steps (about 1 day per alcohol grade) and two full days in histosol for clearing. The tissues were then immersed in hot paraffin (McCormick Scientific Paraplast Plus) in a vacuum oven for 2 days before preparing the blocks and sectioning at 10–12 µm thickness. Sections were rehydrated in histosol and a graded series of ethanol washes (histosol, 100, 90, 70, 50 and 25% ethanol washes in water) and then equilibrated in PBS (Dulbecco’s) before blocking in PBS containing 10% expired FBS and 1% Triton X-100 for 1 h. Primary antibodies [rabbit polyclonal anti-myelin Po and anti-myelin basic protein; Gould, 1992] were added in a 1:500 dilution in PBS and slides were incubated overnight at 4 °C. After washing the sections in PBS for at least 20 min, secondary antibodies (Alexa fluoroprobes conjugated to anti-rabbit or anti-mouse IgG, Invitrogen) were added for 30 min and washed in PBS for immuno-fluorescence visualization and coverslipped with Permount. Pictures of sections were taken using Axiovision LE software (Zeiss TM ) with an AxioCam black and white camera attached to a Zeiss AxioimagerA1 upright fluorescent microscope and assembled into figures 7 and 8 using Adobe Photoshop 7.

Results

Shark Po Antibodies Recognize Different Isoforms of Po

When the antibody that recognizes the full-length Po (RabA in fig. 2) was used in a Western blot of purified shark myelin, two major protein bands were observed. These bands match the molecular weights of the two Po isoforms (28 and 32 kD) previously reported for shark [Saavedra et al., 1989]. A smaller band of lower-molecular-weight (25 kD) protein, not strongly stained, was also apparent. This band has not been previously reported for shark Po.

Immuno-detection using PoCy1 in a sample from the same purified shark myelin revealed three prominent bands at apparent molecular weights of 25, 28 and 32 kD (fig. 2). This peptide antibody, specific for the cytoplasmic domain of shark Po, recognized the same two proteins previously observed with an antibody that recognizes full-length Po as well as a third isoform that matches the smaller band seen in that same Western blot. A Western blot with the PoEx antibody, raised against the N-terminus sequence of shark Po, showed only the two major 28 and 32 kD protein bands. The lack of detection of the smaller 25 kD band with this antibody suggests that this Po isoform is missing at least part of the corresponding N-terminus peptide sequence from its extracellular domain.

The Western blot with PoCy2 antibody detected only the major band at 32 kD (fig. 2). This peptide antibody corresponds to the C-terminus of rat Po, and is 65% homologous to the shark Po C-terminus. The fact that PoCy2 did not detect the 25 and 28 kD protein suggests that these two protein bands are different isoforms of shark Po that lack this end terminus region.

Our finding of several isoforms in the CNS from shark with different truncations has several explanations: that there has been some proteolytic degradation during sample preparation; or that they are due to endogenous proteolytic intermediates present in the lysosomes. The pos-
sibility that protein degradation took place during the myelin preparation and that it can account for the presence of several protein isoforms with decreasing molecular weights can be disregarded because all the procedures were performed with freshly dissected tissues and these were kept at 4°C and in the presence of suitable amounts of protease inhibitors. Second, we did not see any changes in the molecular weight of the major band or an enrichment of the lower band when the lysates were purposefully left overnight at room temperature before running them on a gel (data not shown). Third, these gel patterns have been consistent and reproducible under many experimental conditions, no matter which method of sample preparation (to corroborate the uniqueness of these isoforms, we also ran the proteins under non-denaturing conditions, without β-mercaptoethanol) or the antibody used in the Western blot. The second possibility was that the Po isoforms found in shark are proteolytic intermediates present in the lysosomes. First, if that were the case, we should have observed a more varied number of smaller isoforms in our Western blots, which we did not. Second, we should have also observed varying and/or different amounts of proteins for all the isoforms; but we found almost equal and reproducible amounts of the full length (32 kD) as well of the intermediate isoforms (28 and 25 kD) throughout our Western blots.

**Glycosylation of Shark Po Isoforms**

The myelin Po protein is known to be heavily glycosylated [Uyemura et al., 1981], our finding of three different shark Po protein bands suggests that these might be due to different levels of glycosylation. To determine if this might be the case, we performed an Endoglycase F treatment on the myelin fraction to remove all the sugar residues from shark’s myelin. A Western blot of this fraction showed that these 3 isoforms are fully glycosylated (fig. 3), as each band decreased equally by approximately 6% of their apparent molecular weight after deglycosylation, instead of becoming one single band of 25 kD. This finding suggests that the three different isoforms we observed might be due to products of different Po variants rather than differences in glycosylation.

**Regional Distribution of Po Isoforms**

The regional distribution of these Po isoforms in adult shark nervous system was studied similarly by doing Western blots from solubilized telencephalon, diencephalon, optic tectum, cerebellum and medulla. We found that these CNS regions share the three major isoforms (25, 28 and 32 kD) also recognized by PoCy1 in total purified myelin (fig. 4). Using PoEx antibody showed only the two major bands in these samples, as with total purified shark myelin observed in figure 2 (data not shown). The telencephalon showed the lowest levels of Po, with faint amounts of the 25 kD isoform, whereas cerebellum

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**Fig. 3.** Endo F treatment of myelin Po. Purified shark myelin was treated with an enzyme (EndoF) to remove all carbohydrates. Western blot showed that enzyme treated Po(+) isoforms run about 6% lower than fully glycosylated Po(–). A total of 4 μg of protein from purified myelin was loaded on each lane.

**Fig. 4.** Po isoform distribution in shark brain. Western blots with PoCy1 from different brain regions showed that there was an unequal distribution of shark Po in different areas; 8 μg of total brain lysates was prepared from dissected areas and ran on a SDS-PAGE gel. Tel = Telencephalon; DI = diencephalon; OT = optic tectum; TN = trigeminal nerve; Cer = cerebellum; Med = medulla; SSM = spinal cord myelin.
and diencephalon showed the highest levels of all three isoforms (~50% higher than telencephalon).

We also looked at Po protein in shark PNS using the trigeminal nerve. We found that the expression of Po in the PNS is different from that in CNS. As shown in figure 5, trigeminal nerve has a marked 32 kD band and a second band running at ~27 kD. In order to determine if it was due to different levels of glycosylation, we repeated the process of removing all the carbohydrates from the sample with EndoF. We observed that as expected for a fully glycosylated Po, the band ran 6% lower after EndoF treatment although still at a clearly different weight compared with our standard myelin from spinal cord (fig. 5).

**Developmental Expression of Shark’s Po Isoforms**

Finally, in order to find out if these distinct Po isoforms are regulated or equally expressed throughout development, whole brain from sharks at different stages of development were analyzed. Figure 6 shows Western blots of Squalus acanthias shark at embryonic stages of 4, 4.4, 9 and 22 cm. Levels of Po glycoprotein were detectable only in the 22 cm embryos. We were able to observe only the 28 and 32 kD isoforms, indicating that the 25 kD isoform is not present during the initial stages of shark myelin formation. After over-exposure using ECL detection methods (Enhanced Chemiluminescence) we were able to observe that the 9 cm embryo had low levels of the 28 and 32 kD. A similar finding of fewer isoforms during development has been reported for trout larvae [Jeserich et al., 1990; Stratmann and Jeserich, 1995]. In addition, we observed a 34 kD isoform in the 22 cm embryo (arrow in fig. 6), not present in adult shark brain after extensive over-exposures using Enhanced Chemiluminescence’s detection (data not shown).

We further corroborated that Po is not found in the very early stages of shark development (<5 cm) by performing immunohistochemistry of shark embryo sections (embryos at 5 cm of a different species: Chiloscyllium punctatum) using a RatPo antibody that is capable of detecting Po in fixed tissue (fig. 7A, D, E; data for younger embryos, smaller than 5 cm, not shown). The earliest time that we were able to immunostain for Po was in embryos at 5 cm stage) and onwards [Ballard, 1993]. In the youngest embryos we observed very few myelinated tracts, but in older embryos (6 cm) there were more Po-positive fibers running laterally in the midbrain (fig. 7A, D, E). Meanwhile in the midbrain area there was robust and widespread neurofilament expression, this is a marker for mature neurons (fig. 7B). Because the staining for Po was not striking in these younger embryos, we performed immunostaining for myelin basic protein (MBP) another marker for myelination (fig. 7C). We noticed that MBP expression was detectable at earlier stages and in...
more fibers compared with Po in the *Chiloscyllium* embryos (compare fig. 8C with D). We were able to detect smaller amounts of Po and larger amounts of MBP in stage 32 spinal cords and hindbrain (fig. 8D and A), suggesting that MBP expression might occur before Po. Later on in development (7 cm embryos) we observed robust and comparable amounts of Po and MBP expression in the spinal cords of the embryos as well as in the PNS: ciliary ganglion, spinal nerves and sensory ganglions (fig. 8B, D, E, F).

**Discussion**

Research on non-mammalian myelin Po expression suggested the existence of two to four Po-like glycoproteins in shark and fish CNS [Tai and Smith, 1983, 1984; Saavedra et al., 1989]. Our study demonstrates for the first time that those shark glycoproteins, sometimes referred to as IP1-IP4 [Jeserich and Waehneldt, 1986], are indeed Po isoforms and that there is an additional isoform that was not considered Po in the past. We were able to establish this fact by the use of specific antibodies against shark’s Po peptide sequences.
Previous studies showed that there were two Po isoforms in another elasmobranch species, the gummy shark (*Mustelus antarcticus*), but they did not identify the smallest, 25 kD glycoprotein in whole brain extracts (likely IP4) as Po [Tai and Smith, 1984]. This discrepancy might be explained by: (a) the fact that we observed that this isoform is present in myelin at much lower amounts (~50%) than the other two; (b) that Tai and Smith did not use antibodies for the detection of these Po proteins in their blots and; (c) that they studied a carcharhiniform shark, which has a brain type different from the shark species we used, a squalomorph, that has a simpler laminar brain type. Although we used different shark species for immunohistochemical detection of Po and MBP during development, there is published and unpublished data with *Mustelus canis*, which indicates that in terms of...
expression of these proteins, the results are the same as the ones presented here [Gould, 1992; Gould et al., 1995]. Furthermore, *Squalus* (which we used for the Western blot analysis) is considered a primitive shark and *Chiloscyllium* (which we used for immunohistochemistry) is considered a more modern one. However, looking at their brains, the gross morphology is very different, e.g., the cerebellum of *Squalus* is smooth and *Mustelus* (a species more like *Chiloscyllium*) has a very convoluted brain. In spite of these overall differences, the cellular structures are quite similar and when we used *Mustelus* myelin, we got similar labeling.

The Po isoforms we detected in shark (25, 27, 28 and 32 kD) are fully glycosylated, as shown by their shift (~6%) in mobility in SDS gels, and are differentially expressed throughout the shark’s nervous system. We found that samples from several CNS regions (telencephalon, diencephalon, optic tectum, cerebellum, medulla and spinal cord) all share the same three Po isoforms, whereas the PNS (trigeminal nerve) contains only the 27 and 32 kD. This two PNS bands are not the product of different degrees of glycosylation but rather suggests that at least in trigeminal nerve there may be a separate, distinct isoform of slightly lower molecular weight (27 kD) than the second one (28 kD) widely present in the CNS.

To date, only one Po mRNA transcript from shark was isolated from a cDNA library [Saavedra et al., 1989]. Our finding of several isoforms in the CNS and PNS from shark with different truncations has several explanations: that either these isoforms are due to differential splicing of Po mRNA transcripts; or that these isoforms are due to posttranslational modification of the full-length shark Po. Up until now the best characterized of the myelin Po proteins, besides mammalian, is the one in fish. In the trout researchers found only one mRNA [Stratmann and Jeserich, 1995]. At least two isoforms have been reported for another shark, though these reported isoforms do not have the same molecular weight as the ones reported in this paper [Tai and Smith, 1983, 1984; Zand et al., 1991]. In other words, elasmobranchs seem to have a series of proteolytically degraded forms of myelin Po that fish or mammals do not present.

All these results lead to the suggestion that Po undergoes proteolytical modification. This event is not uncommon and happens to be an effective mechanism of physiological regulation [Neurath, 1989]. It has been observed to occur at the cytoplasmic as well as extracellular N-terminus of several adhesion molecules and receptors [Ozawa and Kemler, 1990; Covault et al., 1991; O’Bryan et al., 1995]. E-Cadherin adhesiveness requires a precise endogenous cleavage of a peptide at its N-terminus domain [Ozawa and Kemler, 1990]. Slit2 is a well known chemorepellent molecule that is also proteolytically cleaved in the cell [Nguyen Ba-Charvet et al., 2001]. Moreover, there is evidence in sharks of post-translational modification of MBP [Deibler et al., 1975; Zand et al., 2001]. Thus, it is quite feasible that during myelination, different proteases might be activated that will modify the expression of Po as it happens for MBP. However, there is as yet no strong evidence towards a splice hypothesis or a proteolytic one.

The PNS and CNS myelin organization of cartilaginous and higher vertebrates depends on the same myelinating glial cells: oligodendrocytes for CNS and Schwann cells for the PNS [Bakay and Lee, 1966; Long et al., 1968; Schweigreiter et al., 2006]. However, the differentiated oligodendrocytes of mammals do not express Po at all but rather the PLP/DM20 glycoproteins [Milner et al., 1985]. Cartilaginous and bony fish have PLP/DM20-like proteins [Kitagawa et al., 1993], but these proteins do not seem to play the role in myelin compaction that is observed in terrestrial vertebrates such as mammals [Boisson et al., 1995; Yoshida and Colman, 1996]. Due to the high levels of Po in both CNS and PNS of non-mammalian vertebrates, it has been hypothesized that Po is responsible for myelin compaction in these organisms. However, at some point during evolution, between cartilaginous fish and reptiles, Po stopped being expressed in CNS, and PLP/DM20 came to play a major role in the myelin formation of higher vertebrates.

Our observations of several isoforms of Po with different truncations could shed some light on the change of role of Po, as well the ‘phenotypically silent drop-out of Po from the terrestrial vertebrate CNS’ [Yoshida and Colman, 1996]. Throughout evolution, protein modifications of Po in CNS myelin, such as the proteolytically cleaved products we present in this paper, could have made it less capable of sustaining compact myelin, allowing the adhesive PLP/DM20-like glycoproteins, already present as potential adhesive proteins, to take over that role in myelogenesis. Wong and Filbin have already shown that a truncated rat Po at its cytoplasmic terminus is incapable of supporting homophilic adhesion of transfected CHO cells and that, furthermore, it has a dominant negative effect on the adhesion of the full-length Po [Wong and Filbin, 1996]. The isoform of Po with such an effect is missing the same sequence recognized by the PoCy2 antibody. This sequence is absent as well in the 28, 27 and 25 kD isoforms in shark’s Po. Maybe the post-
translational modifications of full-length Po in elasmobranches stripped it of its adhesive properties, leaving an open door for the PLP/DM20 protein to substitute for Po as the major adhesive molecule in the CNS. More than a ‘silent drop’ of Po as the major adhesive component in CNS myelin, it was a ‘takeover’ by the PLP/DM-20 molecules. It remains to be seen whether each of the identified isoforms of shark Po can support homophilic adhesion. The fact that this protein is so highly conserved throughout evolution suggests that those forms lacking the cytoplasmic domain might not support homophilic adhesion, or at least not in the same fashion as the full-length protein.

The pattern for Po shown in this paper is very different from the one found in higher vertebrates, where Po is present only in the PNS, and only as a single isoform with slight variations in its levels of glycosylation. The current result of several Po isoforms, spatially and developmentally regulated, is a novel finding for this major myelin glycoprotein.

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