DSD-1-Proteoglycan Is the Mouse Homolog of Phosphacan and Displays Opposing Effects on Neurite Outgrowth Dependent on Neuronal Lineage

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DSD-1-PG is a chondroitin sulfate proteoglycan (CSPG) expressed by glial cells that can promote neurite outgrowth from rat embryonic mesencephalic (E14) and hippocampal (E18) neurons, an activity that is associated with the CS glycosaminoglycans (GAGs). Further characterization of DSD-1-PG has included sequencing of peptides from the core protein and the cloning of the corresponding cDNA using polyclonal antisera against DSD-1-PG to screen phage expression libraries. On the basis of these studies we have identified DSD-1-PG as the mouse homolog of phosphacan, a neural rat CSPG. Monoclonal antibodies 3H1 and 3F8 against carbohydrate residues on rat phosphacan recognize these epitopes on DSD-1-PG. The epitopes of the antibodies, L2/HNK-1 and L5/Lewis-X, which have been implicated in functional interactions, are also found on DSD-1-PG. Although DSD-1-PG has previously been shown to promote neurite outgrowth, its upregulation after stab wounding of the CNS and its localization in regions that are considered boundaries to axonal extension suggested that it may also have inhibitory functions. Neonatal dorsal root ganglion (DRG) explants grown on a rich supportive substrate (laminin) with and without DSD-1-PG were strikingly inhibited by the proteoglycan. The inhibitory effects of DSD-1-PG on the DRG explants were not relieved by removal of the CS GAGs, indicating that this activity is associated with the core glycoprotein. The neurite outgrowth from embryonic hippocampal neurons on laminin was not affected by the addition of DSD-1-PG. This indicates that DSD-1-PG/mouse phosphacan can have opposing effects on the process of neurite outgrowth dependent on neuronal lineage.

Key words: DSD-1-proteoglycan; phosphacan; chondroitin sulfate proteoglycan; neurite outgrowth; hippocampus; dorsal root ganglion

Extracellular matrix (ECM) molecules are expressed in many regions of the developing nervous system where they are thought to play crucial roles in neural development, being implicated in processes such as neuronal adhesion, shape, migration, and neurite outgrowth (Lander and Calof, 1993; Letourneau et al., 1994; Faissner and Steindler, 1995; Fitch and Silver, 1997). Although the organization of the ECM in the vertebrate CNS is not well understood it is marked by the relative abundance of chondroitin sulfate proteoglycans (CSPGs) and hyaluronan (Maleski and Hockfield, 1997; Rauch, 1997).

CSPGs, a heterogeneous set of proteins bearing glycosaminoglycans (GAGs) of the chondroitin sulfate (CS) class, account for most of the “soluble” proteoglycans in the brain (Lander, 1993). During development, strong immunostaining for CS often localizes to territories thought to act as barriers to migrating neurons or extending axons such as the roof plate and midline dorsal tectum (Snow et al., 1990b; Katoh-Semba et al., 1995), the posterior sclerotome (Oakley and Tosney, 1991; Landholt et al., 1995), and the dorsal root entry zone and dorsal columns in the spinal cord (Pindzola et al., 1993). In vitro CSPGs (Snow et al., 1990a; Dou and Levine, 1994; Maeda and Noda, 1996), the isolated core proteins of CSPGs (Dou and Levine, 1994; Maeda and Noda, 1996), and CS by itself (Snow et al., 1990a; Smith-Thomas et al., 1994) have also been shown to inhibit cell migration or neurite outgrowth on defined growth-promoting substrata.

Yet, it is clear that tissues that express CS do not always exclude the entry of axons, and in some cases CS staining coincides with developing axon pathways (Sheppard et al., 1991; Bicknese et al., 1994; McAdams and McLoon, 1995). Indeed, several in vitro studies suggest that CSPGs (Streit et al., 1993; Faissner et al., 1994), CS (Lafont et al., 1992; Fernaud-Espinosa et al., 1994), and isolated core proteins (Iijima et al., 1991) can promote rather than inhibit neurite outgrowth.

The mouse neural CSPG, DSD-1-PG, was originally identified by the monoclonal antibody (mAb) 473HD, which specifically recognizes the CS epitope DSD-1 (Faissner, 1988; Faissner et al., 1994).
1994), DSD-1-PG promotes neurite outgrowth from rat embryonic mesencephalic (E14) and hippocampal (E18) neurons, a process that is dependent on the DSD-1 GAG epitope, because it can be blocked specifically by the mAb 473HD or by enzymatic digestion of the CS GAGs (Faisser et al., 1994). The DSD-1 epitope has been shown to be sulfation-dependent and to correlate with CS D motifs (Clement et al., 1998). In cerebellar cultures, it is expressed on the surface of immature glial cells where it is downregulated with ongoing maturation, and it is not expressed on neurons (Schnädelbach et al., 1998). The developmental distribution of DSD-1-PG has been found to correspond to regions related to the formation of axonal trajectories such as the interrhombic boundaries in chick, which is consistent with a role in growth promotion (Heyman et al., 1995), but it has also been found at glial barrel field boundaries in the developing mouse somatosensory cortex, which neurons do not cross (Stein­dler et al., 1995).

Several studies have reported the enhanced expression of growth-inhibiting CSPGs in the context of CNS lesions (McKeon et al., 1991; Bovolenta et al., 1993; Pindzola et al., 1993; Lips et al., 1995), and DSD-1-PG is also strongly upregulated after wounding in the CNS (Laywell et al., 1990; Laywell and Stein­dler, 1991; Barker et al., 1996; Deller et al., 1997). Such an upregulation of DSD-1-PG in the wound reaction might be attributable to the action of TGF-β based on studies of Oli-neu, an oligodendrocyte precursor cell line (Schnädelbach et al., 1998).

Additional in vitro studies of the effects of DSD-1-PG on neurite outgrowth have been made, and we report here an inhibitory activity of DSD-1-PG on dorsal root ganglion (DRG) explants. Further biochemical characterization of DSD-1-PG is also presented, together with the cloning of the cDNA corresponding to the core protein and its identification as the mouse homolog of phosphacan.

**MATERIALS AND METHODS**

*Antibodies and immunohistochemistry.* Rat mAb 473HD and the rabbit polyclonal antibody (pAb) KAF13 were described previously (Faisser, 1988; Faisser et al., 1994). 473HD, a rat mAb, should not be confused with another mAb called 473, from mouse, which also recognizes a CS epitope on a rat neural CSPG (Kosaka et al., 1992). mAbs 3F8, 3H1, and 2B4 against phosphacan and phosphacan-KS are available from the Developmental Studies Hybridoma Bank (University of Iowa, Department of Biological Sciences, Iowa City, IA). mAbs L2/4HK-1 (clones 334, rat IgM), L3 (clone 492, rat IgM), L4 (clone 483, rat IgM), and LS (clone 487, rat IgM) have been described previously (Kruse et al., 1984; Kücherer et al., 1987; Streit et al., 1990). Western blots were performed as described and developed with peroxidase-derivatized secondary antibodies using the ECL kit (Amersham Buchler GmbH, Braunschweig, Germany) according to the manufacturer’s instructions. Iodination and immunoprecipitation of DSD-1-PG were performed as described previously (Faisser et al., 1994).

*Protein purification and deglycosylation studies.* The DSD-1-PG was purified from detergent-free physiological saline-buffered brain lysates from postnatal day (P) 7–14 mice as described previously using a combination of affinity chromatography with the mAb 473HD bound to Sepharose resin, and anion-exchange chromatography (Faisser et al., 1994). It was quantitated using the protein assay (Bio-Rad Laboratories, Munich, Germany) or the Micro BCA reagent (Pierce, Rockford, IL) and by the determination of uronic acid equivalents (Blumenkrantz and Asboe-Hansen, 1973). The DSD-1-PG was deglycosylated in the presence of protease inhibitors (Faisser et al., 1994) using chondroitinase ABC (ChABC) (EC 4.2.2.4; Boehringer Mannheim GmbH, Mannheim, Germany) at 50 mU/ml in 40 mM Tris, pH 8.4, 40 mM sodium acetate, 0.01% BSA for 2 hr at 37°C, and keratanase (EC 3.2.1.103; Seikagaku Kogyo, Tokyo) at 100 mU/ml in 50 mM Tris, pH 7.4, for 2 hr at 37°C.

*Protein sequencing.* Purified DSD-1-PG was digested with ChABC and run out on a 4–10% SDS-PAGE. Because of the poor staining of DSD-1-PG, the Coomassie brilliant blue band corresponding to the core glycoprotein was excised based on alignments with adjacent lanes that were either silver-stained or blotted with KAF13. The gel band was then rinsed in water to remove acetic acid, crushed, and incubated overnight at 37°C in 0.1 M sodium hydrogen carbonate, pH 8.3, with either 1 U sequencing grade trypsin (Boehringer Mannheim) or subtilisin (Sigma, Munich, Germany) at 37°C for 24 hr. The resulting peptides were recovered from the gel pieces by extraction with acetonitrile and concentrated in a lyophylizer. The peptides were then loaded on a reverse-phase HPLC (RP-HPLC) nanocolumn in 1% aqueous trifluoroacetic acid and 0.1% TFA from 10 to 80% acetonitrile gradient. This yielded a number of peptides from which the N-terminal amino acid sequence was obtained by Edman degradation using a protein sequencer (Applied Biosystems Sequencer 477A/494; Applied BioSystems, Foster City, CA).

**Screening of cdna expression libraries.** Several mouse brain λ cDNA expression libraries were screened using the pAb KAF13 at 1 μg/ml, following standard procedures. The libraries used were BALB/c neona­tal whole-brain oligo-dT and random-prime Uni-ZAP XR 1 (Strategene GmbH, Heidelberg, Germany), C57/B16 neonatal and adult whole-brain oligo-dT-primed λ g11, and C57/B16 neonatal cerebellum (P2–6) oligo-dT-primed λ g11 (Barthels et al., 1988). The DNA sequence of the positive clones was obtained using the Cy5 AutoRead screening kit and the AL Fexpress DNA sequencing system (Pharmacia Biotech, Freiburg, Germany). After screening, more than 800 positive clones were obtained: TB24 and p159, which correspond to bases 1144–1509 and 1032–3591, respectively, of the cdna sequence shown in Figure 2, and p166, which is the short, spliced form of receptor protein tyrosine phosphatase (RPTP)β/c with bases 1186–2392 of the phosphacan sequence, the splice site, and then bases 4951–7850 of the rat RPTPβ/c cdna sequence (RN09357 in the EMBL database) corre­sponding to the entire transmembrane domain, two tyrosine phosphatase domains, and 180 bases of the 3′ untranslated region of the cdna sequence. A Marathon cdna amplification kit (Clontech, Heidelberg, Germany) was used to generate a P7 cerebellum cdna library from 1 μg poly(A+) RNA. The 5′ end of the sequence was obtained from this cdna library using RACE (rapid amplification of cdna ends) PCRs with specific primers based on the sequence of clone p159, and 5′-priming from an adaptor sequence ligated to the cdna. The PCR products were subcloned into plasmid pCR 2.1 (Invitrogen, Leck, The Netherlands) and sequenced. Once the 5′ end was known, the sequence was verified by PCR amplification using specific 5′ primers and sequenc­ing the corresponding PCR products. At the 3′ end, an expressed sequence tag (EST) clone, MM1180153 (EMBL database), was identified by sequence comparison of the rat sequence with computer DNA data­bases. This EST clone was obtained from the IMAGE consortium (RZPD, Berlin, Germany) and on sequencing was found to correspond to bases 1144–1509 and 1032–3591 of the cdna sequence shown in Figure 2. This clone and clone p159 were used to generate the intervening 3′ sequence (3510–5064) from the cdna library using PCR. The sizes of mRNAs corresponding to the clone p159 were investigated by Northern bloting using P7 cerebellum poly(A+) RNA and the Church method.

**Cell culture.** Hippocampal cell cultures were established from embryonic day 18 (E18) rat brains as described previously (Faisser et al., 1994; Kempl et al., 1996). Briefly, the hippocampal neurons were obtained by microdis­section and dissociated in supplemented N2 medium (Banker and Cowan, 1977). The resulting cell suspension was plated on coverslips at a density of 10,000 cells/cm² in supplemented N2 medium. DRG explants from P0 or P1 mice were prepared as described previously (Götz et al., 1996). Briefly, the ganglia were placed in HBSS on ice, and tissue and nerve fibers sticking to the explants were removed (Seilheimer and Schachner, 1988). Whole ganglia were transferred into culture medium consisting of DMEM medium supplemented with 15 ng/ml nerve growth factor (Sigma), and 20 μM arabinosylcytosine (Sigma). These two neurite outgrowth model systems were chosen for analysis because they have been well characterized and standardized in our laboratory for a range of conditions. Although the hippocampal neurons are from rat and the DRG explants are from mouse, the interspecies differences are not very pronounced between rat and mouse. For these reasons it seemed adequate to use them for studies of the effects of DSD-1-PG as a substrate on poly-ornithine and laminin, respectively.

There is in fact an extensive literature demonstrating the wide-ranging cross-species effects of extracellular matrix components, for example concerning laminin-1 or fibronectin, which are isolated from outdated human plasma or Engelbreth-Holm-Swarm-tumor tissue, respectively.
and nevertheless used for neurons from various species, with comparable results.

Neurite outgrowth assays. Neurite outgrowth of E18 hippocampal neu-
rons plated on supports was conducted as described previously (Locher et al., 1991; Faissner et al., 1994; Götz et al., 1996). In brief, glass
coverslips were treated with 1.5 μg/ml poly-bi-ornithine (PORN; Sigma) in
0.1 M borate buffer, pH 8.2, for 1 hr at 37°C in a humidified atmo-
sphere. Laminin (laminin-1 from Engelbreth-Holm-Swarm mouse sar-
coma cells; Boehringer Mannheim) was subsequently coated at 5 μg/ml
in 50 μl PBS per coverslip for at least 1 hr at 37°C in the incubator, and
the coverslips were washed twice with PBS before E18 hippocampal
neurons were plated at 10,000 cells/cm². DSD-1-PG was coated at 5 μg
uronic acid equivalents per milliliter. After coating, the Petri dishes were
washed three times with PBS and blocked by incubation with 0.25% BSA
for 2 hr. After 24 hr of culture, neurons were stained for tubulin using an
mAb to α-tubulin (clone DM 1A; Sigma) as described previously (Götz et al., 1996). The quantitative morphometry and statistical analysis of
neurite outgrowth were as described previously. Neurite outgrowth was
determined as the fraction of process-bearing cells from at least 100
neurons per coverslip chosen at random and given in percentage. The
morphometric analysis of neurite lengths was performed with the system
Quantimet 500 MC (Leica, Bensheim, Germany) by measuring neurons
with a process longer than one neuronal cell body. At least three
independent experiments were analyzed, and the data were evaluated
using nonparametric statistics. The distribution of the longest neurites
was compared using the Mann–Whitney U test; the mean values of sums
of the longest neurites were compared by t test. P0/P1 mouse DRG
explants were grown in tissue culture plastic Petri dishes (3 cm diameter)
(Nunc, Wiesbaden, Germany) coated with PORN, laminin, and purified
murine collagen (Nunc) (Faissner et al., 1994) were obtained from proteolytic fragments of the
protein sequence of the secreted rat CSPG phosphacan (Maurel et al., 1994). As illustrated in Figure 3, there is
a predicted signal peptide in the first 24 residues, followed by a
carbonic anhydrase-like domain from 34 to 302 and a fibronectin
carboxyterminal domain from 312 to 406, although the remainder of the
protein has no apparent strong homology with other known
proteins. The relationship of the mouse RPTPζ/β to the DSD-1-PG occurred as a very high molecular mass smear that has
been estimated by gel filtration to be ~800–1000 kDa (Faissner et al., 1994). This is first detected around E13 and then shows a

RESULTS

Peptide sequences from DSD-1-PG correspond to phosphacan, and screenings of brain cDNA expression
libraries with DSD-1-PG antiserum yield phosphacan clones

Further characterization of the primary structure of DSD-1-PG
entailed the obtention of peptide sequences from the core pro-
tein, together with the screening of mouse brain cDNA expres-
sion libraries using a polyclonal antiserum raised against
DSD-1-PG.

A large number of peptide sequences corresponding to the
protein sequence of the secreted rat CSPG phosphacan (Maurel et al., 1994) were obtained from proteolytic fragments of the
purified DSD-1-PG (Fig. 1), suggesting that DSD-1-PG may be
the mouse equivalent of this protein. This observation was sup-
ported by our screening of several bacterially expressed mouse-
brain cDNA phage libraries using the pAb KAF13, which was
previously raised against the purified DSD-1-PG (Faissner et al., 1994). On Western blots, the antiserum preferentially recognizes
the ChABC-digested core protein rather than whole PG (Fig. 4),
and hence during screening it should recognize protein epitopes
corresponding to the protein core. The three positive clones that
we obtained on stringent screening of 3

PCR amplification. This includes the entire 4.8 kb coding se-
quence and the corresponding 1612 amino acid sequence of
the protein.

Overall, the mouse homolog has 90% identity (97% similarity
based on semi-conservative amino acid substitutions) with the rat
sequence (Maurel et al., 1994). As illustrated in Figure 3, there is
a predicted signal peptide in the first 24 residues, followed by a
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As further evidence of the identity of DSD-1-PG as mouse phosphacan, the developmental expression profiles of the proteo-
glycans were compared, as was the cross-reactivity of various
antibodies. In Western blots of mouse-brain extracts (Fig. 4),
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Peptide sequences

Trypsin digest

Garwood et al. • Molecular Characterization of DSD-1-PG

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The 6.8 kb nucleotide sequence of the DSD-1-PG/mouse phosphacan cDNA is shown together with the deduced 1612 amino acid sequence of the 4.8 kb open reading frame. 1 marks the start of the coding region; * indicates the stop codon at the end. The 3' untranslated region ends in the poly(A) tail. The accession number for this sequence in the EMBL database is AJ133130.


Figure 2. Nucleotide sequence of DSD-1-PG/mouse phosphacan. The 6.8 kb nucleotide sequence of the DSD-1-PG/mouse phosphacan cDNA is shown together with the deduced 1612 amino acid sequence of the 4.8 kb open reading frame; +1 marks the start of the coding region; * indicates the stop codon at the end. The 3' untranslated region ends in the poly(A) tail. The accession number for this sequence in the EMBL database is AJ133130.
dramatic increase in expression to plateau around the second postnatal week, although it remains at high levels in the adult brain (Fig. 4a). When the CS GAGs of DSD-1-PG are digested by chondroitinase ABC, the CS epitope recognized by mAb 473HD is also removed. The resulting core glycoprotein, which can be visualized with the pAb KAF13, has a relative molecular mass of ~350-400 kDa and shows the same developmental profile as the whole CSPG (Fig. 4b). This expression pattern is very similar to that given for rat phosphacan (Meyer-Puttlitz et al., 1995).

DSD-1-PG can be readily extracted from CNS tissues using mild buffers in the absence of detergent (Faissner et al., 1994). As the mouse homolog of phosphacan, and hence the secreted form of RPTPβ, antibodies recognizing DSD-1-PG should also recognize the receptor isoforms. In membrane preparations extracted with 1% w/v NP40-containing buffers, it was indeed possible to visualize bands on Western blots corresponding to the transmembrane RPTPβ isoforms with 473HD and KAF13 (data not shown). The high molecular weight smear in the extracts could be digested with chondroitinase ABC to give core glycoproteins ~400 and 250 kDa (the long and short RPTPβ, respectively), similar to the pattern in rat tissue (Sakurai et al., 1996).

From the cDNA sequence, the predicted size of the core protein of DSD-1-PG/mouse phosphacan is 175 kDa. The discrepancy between the theoretical mass and the observed migration of the core protein after ChABC digestion is an indication of the degree of other carbohydrate modifications on the protein. There are 16 potential N-glycosylation sites in both rat and mouse phosphacan [PROSITE; Bairoch et al. (1997)] and approximately 30 predicted O-glycosylation sites (Hansen et al., 1995) (Fig. 3b).

In rat phosphacan, a distinction has been made based on mutually exclusive carbohydrate epitopes recognized by the mAbs 3F8 and 3H1 (Rauch et al., 1991; Maurel et al., 1994). The 3F8 N-linked oligosaccharide epitope is present on phosphacan, whereas 3H1 reacts with a keratan sulfate epitope present on a keratan sulfate glycoform of phosphacan, termed phosphacan-KS (Maurel et al., 1994). As can be seen in Figure 5, the purified DSD-1-PG is recognized by both 3H1 and 3F8, indicating the presence of both epitopes, and the 3H1 reactivity on DSD-1-PG can be digested by keratanase. Because DSD-1-PG is purified by affinity chromatography with the 473HD mAb, the implication is that the DSD-1 epitope is present, at least in mouse, on both the phosphacan and phosphacan-KS glycoforms. A protein epitope in the N-terminal half of rat phosphacan recognized by another mAb, 2B49 (Sakurai et al., 1996), is apparently absent from DSD-1-PG (data not shown).

Two other characterized carbohydrate epitopes recognized by mAbs L2 and L5 have also been found on DSD-1-PG (Fig. 6).
recognizes the HNK-1 epitope, a sulfated glucoronic acid carbohydrate found on several neural recognition molecules, such as L1 and N-CAM (Kruse et al., 1985), which has been shown to be involved in the migration of neural crest cells in vivo (Bronner-Fraser, 1987) and has been implicated in vitro in neuronal migration, neurite outgrowth (Lallier and Bronner-Fraser, 1991; Lallier et al., 1992), and short-term cell-to-cell and cell-to-laminin adhesion (Keilhauer et al., 1985; Kunemund et al., 1988; Hall et al., 1993). It was not possible to remove the L2/HNK-1 epitope from DSD-1-PG using peptide-N-glycosidase F, keratanase, or ChABC (data not shown), an observation similar to that found with rat phosphacan (Rauch et al., 1991; Maeda et al., 1995).

The L5 mAb has been shown to recognize the Lewis-X N-linked carbohydrate structure (Streit et al., 1996), which appears to play a role in early neurogenesis (Roberts et al., 1991) and astrocytic process formation (Streit et al., 1993). Lewis-X, as recognized by the mAb FORSE-1, has also been found on rat phosphacan (Allendorfer et al., 1995). The L5 antibody has also been used to characterize astrochondrin, a large mouse CSPG expressed by mature astrocytes (Streit et al., 1990, 1993). On digestion of its CS GAGs, astrochondrin migrates as three core proteins at ~380, 360, and 260 kDa. It also carries the L2/HNK-1 epitope and the DSD-1 epitope (Streit et al., 1993) and has been described as the only L5-bearing CSPG expressed by astrocytes, showing a developmental profile similar to DSD-1-PG (Streit et al., 1990, 1993), which is also expressed by GFAP+ astrocytes (Schnädelbach et al., 1998). As such, it is possible that there is some identity between DSD-1-PG and astrochondrin.

Other CSPGs have been described whose core proteins show a relative migration similar to DSD-1-PG on SDS-PAGE, but in the absence of peptide sequences or antibody cross-reactivity further identification would be hazardous.

The epitope recognized by mAb 6B4 has been found on migrating neurons (Maeda et al., 1995), and in situ hybridization analysis of sections from the developing brain indicate that there are subsets of neurons that express RPTPβ and phosphacan (Snyder et al., 1996). In the case of DSD-1-PG, colocalization of several antibody markers on cerebellar cultures in vitro does not show an overlap of L1 as a neuronal marker with DSD-1 (Faißner et al., 1994; Schnädelbach et al., 1998), nor in various immunohistochemical studies has a staining of neurons been evident (Gates et al., 1995; Heyman et al., 1995; Steinandler et al., 1995; Wintergerst et al., 1996), although this may reflect higher levels of DSD-1-PG expression and secretion by glial cells compared with neurons. Perhaps in situ hybridization studies using appropriate cDNA probes will prove more sensitive in this respect.

**DSD-1-PG/mouse phosphacan promotes neurite outgrowth from hippocampal neurons grown on PORN but inhibits neurite outgrowth from DRG explants grown on laminin**

The identification of DSD-1-PG as the mouse equivalent of rat phosphacan presents some contradictions concerning the functional activity of the proteoglycan in neurite outgrowth assays. Although DSD-1-PG has clear outgrowth-promoting properties when tested on rat E14 mesencephalic and E18 hippocampal neurons (Faißner et al., 1994) and phosphacan has also been shown to promote morphological differentiation of E16 rat cortical neurons grown at low density on poly-L-lysine (Maeda et al., 1995), phosphacan (Allendorfer et al., 1995). The L5 antibody has also been recognized by the mouse equivalent of rat phosphacan (Allendorfer et al., 1995). The L5 antibody has also been used to characterize astrochondrin, a large mouse CSPG expressed by mature astrocytes (Streit et al., 1990, 1993). On digestion of its CS GAGs, astrochondrin migrates as three core proteins at ~380, 360, and 260 kDa. It also carries the L2/HNK-1 epitope and the DSD-1 epitope (Streit et al., 1993) and has been described as the only L5-bearing CSPG expressed by astrocytes, showing a developmental profile similar to DSD-1-PG (Streit et al., 1990, 1993), which is also expressed by GFAP+ astrocytes (Schnädelbach et al., 1998). As such, it is possible that there is some identity between DSD-1-PG and astrochondrin.

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Other CSPGs have been described whose core proteins show a relative migration similar to DSD-1-PG on SDS-PAGE, but in the absence of peptide sequences or antibody cross-reactivity further identification would be hazardous.

The epitope recognized by mAb 6B4 has been found on migrating neurons (Maeda et al., 1995), and in situ hybridization analysis of sections from the developing brain indicate that there are subsets of neurons that express RPTPβ and phosphacan (Snyder et al., 1996). In the case of DSD-1-PG, colocalization of several antibody markers on cerebellar cultures in vitro does not show an overlap of L1 as a neuronal marker with DSD-1 (Faißner et al., 1994; Schnädelbach et al., 1998), nor in various immunohistochemical studies has a staining of neurons been evident (Gates et al., 1995; Heyman et al., 1995; Steinandler et al., 1995; Wintergerst et al., 1996), although this may reflect higher levels of DSD-1-PG expression and secretion by glial cells compared with neurons. Perhaps in situ hybridization studies using appropriate cDNA probes will prove more sensitive in this respect.
and CSPGs on DRG explants (Dou and Levine, 1994, 1995; Smith-Thomas et al., 1994; Katoh-Semba et al., 1995; Snow et al., 1996).

As can be seen in Table 1, plating the hippocampal neurons and DRG explants on laminin results in a doubling of both the percentage of neurite-bearing neurons and the mean length of these processes compared with the PORN substrate alone. The neurons on the DSD-1-PG/laminin substrate again responded differently (Table 1; Figs. 8, 9). Although there was no significant change in the outgrowth from the hippocampal neurons, the addition of DSD-1-PG to the laminin substrate resulted in a very striking reduction in both the percentage of neurite-bearing explants and the process lengths for the DRG explants. Thus, it appears that DSD-1-PG can indeed also inhibit neurite outgrowth but that these effects are dependent on the neuron cell type.

The inhibitory effects of CSPGs in some other assay systems have been attributed to the CS GAGs rather than the whole proteoglycan or the core protein (Fernaud-Espinosa et al., 1994; Smith-Thomas et al., 1994; Dou and Levine, 1995). In the case of DSD-1-PG, on the contrary, it is the outgrowth-promoting activity on hippocampal neurons that has previously been shown to be dependent on the CS GAG chains (Faissner et al., 1994; Clement et al., 1998). Indeed, in addition to being removed by ChABC digestion, this is an activity that can be specifically neutralized by the mAb 473HD, implying that the DSD-1 epitope plays a direct role (Faissner et al., 1994). Here, however, removal of the CS GAGs by ChABC digestion of the DSD-1-PG substrate before plating of the neurons had no significant effect on either of the neuronal types studied, there being no alleviation of the inhibition observed with the DRG explants (Table 1). As such, it appears that the observed inhibition of the DRG outgrowth in this assay is associated with the core glycoprotein of DSD-1-PG rather than the CS GAGs.

**DISCUSSION**

We report here the identification of the core protein of DSD-1-PG as the mouse homolog of phosphacan. In *in vitro* studies we have demonstrated that, under the same culture conditions, the purified DSD-1-PG can promote neurite outgrowth from one neuronal type and inhibit it from another.

**Neurite outgrowth and CSPGs**

CSPGs are generally regarded as barriers for neurite outgrowth (Snow et al., 1990a,b; Oohira et al., 1991; Dou and Levine, 1994; Milev et al., 1994). They have been shown to be associated with glial boundaries in the development of many CNS structures, including the roof plate and midline dorsal tectum, posterior sclerotome, dorsal root entry zone, and dorsal columns in the spinal cord, locations that are considered to be axon inhibitory regions (Fitch and Silver, 1997). *In vitro* studies show that CSPGs can inhibit neurite outgrowth and elongation (Snow et al., 1990a; Fichard et al., 1991; Oohira et al., 1991; Brittis et al., 1992; Dou and Levine, 1994; Maeda and Noda, 1996), effects that can be associated with either the whole PG, the CS GAGs (Snow et al., 1990a; Schwab et al., 1993), or the protein cores (Oohira et al., 1991; Grumet et al., 1993).

It is now clear, however, that there are also regions in which CSPGs can inhibit neurite outgrowth and elongation (Snow et al., 1990a; Fichard et al., 1991; Oohira et al., 1991; Brittis et al., 1992; Dou and Levine, 1994; Maeda and Noda, 1996), effects that can be associated with either the whole PG, the CS GAGs (Snow et al., 1990a; Schwab et al., 1993), or the protein cores (Oohira et al., 1991; Grumet et al., 1993).

**Neurite outgrowth and DSD-1-PG/mouse phosphacan**

There has been some confusion about the effects of phosphacan on neurite outgrowth. It has been shown to inhibit neurite growth from E9 chicken brain neurons plated on NgCAM (Grumet et al., 1993; Milev et al., 1994) but to promote morphological differentiation of E16 rat cortical neurons grown at low density on poly-L-lysine (Maeda et al., 1995; Maeda and Noda, 1996). The study with phosphacan plated on poly-L-lysine (Maeda and Noda, 1996) showed that the percentage of neurite-bearing cells was higher for both E16 cortical and E16 thalamic neurons but that although the length of the resulting neurites from cortical neurons was increased relative to poly-L-lysine, there was no corresponding change in the thalamic neurons.

Although these studies were with relatively crude mixtures of many different neuronal subtypes, they are not inconsistent with our contrasting observations of, on the one hand, a cell type-specific neurite outgrowth promoting activity of DSD-1-PG/mouse phosphacan from hippocampal and mesencephalic neurons plated on PORN, yet on the other hand, a striking inhibition by DSD-1-PG of neurite outgrowth from DRG explants plated on laminin.

The inhibitory effects of DSD-1-PG are not removed by ChABC digestion, indicating that they are associated with the core glycoprotein. Interestingly, digestion of the CS GAG chains did not affect either the inhibitory or the promotory effects of phosphacan (Milev et al., 1994; Maeda and Noda, 1996), and although we have also found that the digestion of the CS GAG chains does not alleviate the inhibitory effects of DSD-1-PG on

**Table 1. Summary of neurite outgrowth assays comparing the effect of DSD-1-PG coated on PORN or laminin on hippocampal neurons and DRG explants**

<table>
<thead>
<tr>
<th>Substrate</th>
<th>E18 Hippocampal neurons</th>
<th>P0/P1 DRG explants</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total no. of analyzed</td>
<td>Neurite-bearing</td>
</tr>
<tr>
<td></td>
<td>neurons</td>
<td>cells (%) ±SE</td>
</tr>
<tr>
<td>PORN</td>
<td>300</td>
<td>40.3 ± 3</td>
</tr>
<tr>
<td>PORN + DSD-1-PG</td>
<td>300</td>
<td>62.3 ± 4</td>
</tr>
<tr>
<td>LN</td>
<td>300</td>
<td>79.0 ± 4</td>
</tr>
<tr>
<td>LN + DSD-1-PG</td>
<td>300</td>
<td>87.0 ± 3</td>
</tr>
<tr>
<td>LN + DSD-1-PG/ChABC</td>
<td>300</td>
<td>86.0 ± 2</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>30.0 ± 1</td>
</tr>
<tr>
<td></td>
<td>35</td>
<td>28.6 ± 1</td>
</tr>
<tr>
<td></td>
<td>43</td>
<td>65.1 ± 4</td>
</tr>
<tr>
<td></td>
<td>44</td>
<td>31.8 ± 3</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>35.0 ± 1</td>
</tr>
</tbody>
</table>

Total number of analyzed neurons/DRG explants out of three independent experiments. SE, Standard error (t test). LN, Laminin.

*Data from two experiments.
DSD-1-PG on hippocampal neurons (HCN) and dorsal root ganglion explants (DRG). A, The percentage change in neurite-bearing cells/explants versus the PORN control; B, the percentage of neurite outgrowth versus PORN. In the case of hippocampal neurons the length of the longest neurites was measured, whereas for the DRG explants, the surface area of the halo in square millimeters was measured. At least three independent experiments were analyzed, and the mean values of the neurite lengths/halos and percentage of neurite-bearing cells/explants were compared by t test. n.s., Nonsignificant; "" 0.01 > p > 0.001. Error bars represent the SE.

Bacterially and eukaryotically expressed protein domain constructs corresponding to different parts of the core protein of phosphacan/RPTPβ/ζ have been tested in neurite outgrowth assays (Peles et al., 1995, 1997; Sakurai et al., 1997). These indicated that outgrowth from chick tectal neurons could be supported by the carbonic anhydrase domain, an effect potentiated by addition of the “S” domain (Fig. 3).

In addition to the CS GAGs, DSD-1-PG is highly glycosylated with other carbohydrate modifications, as illustrated by the presence of the 3H1, 3F8, L2/HNK-1, and L5/Lewis-X epitopes. It appears likely that the sulfation, carbohydrate composition, and oligosaccharide structure of DSD-1-PG/phosphacan is developmentally regulated and that at least some of these carbohydrate modifications could alter its affinity for other proteins, for example, the N-linked sugars on the carbonic anhydrase and FN III
domains of phosphacan that mediate its interactions with NgCAM, NCAM, and Tenascin-C (Milev et al., 1995).

In effect, DSD-1-PG/phosphacan has been shown to possess three levels at which it can interact with other molecules either in the ECM or on cell membranes. These are the GAG chains, the other N- and O-linked oligosaccharides, and finally the regions of the protein core that are not covered by carbohydrate modifications. With such a range of possible interactions, it is not surprising that DSD-1-PG/phosphacan is implicated in many developmental processes such as migration and neurite outgrowth. In addition to variations in the presentation of such sites of interaction on the PG, the amplitude of its effects is likely to be dependent on localized combinatorial variations, both quantitative and qualitative, of promoting and inhibitory factors that recognize these sites. Cell type-specific differences in the cell surface receptors when confronted with various potential ligands in the ECM, and the relative responsiveness of their intracellular signaling mechanisms to such factors, could then account for the differential cellular behavior observed.

**In vivo distribution of DSD-1-PG/mouse phosphacan**

The expression profile of DSD-1-PG matches that reported for rat phosphacan, with a rapid increase in its concentration during the late embryonic and early postnatal period, the levels remaining high in adult brain (Meyer-Puttlitz et al., 1995).

The distribution of DSD-1-PG during development has been found to correspond to regions related to the formation of axonal trajectories. In this respect, it might play either a neurite-promoting role, as in the interrhombic boundaries in chick (Heyman et al., 1995), or an inhibitory role, which would correspond to its presence in glial barrel field boundaries in the developing somatosensory cortex of mouse (Steindler et al., 1995).

Studies of the distribution of both the phosphacan mRNA (Engel et al., 1996) and the expressed protein (Meyer-Puttlitz et al., 1996) show that at E13–16, for example, the phosphacan mRNA is largely confined to areas of active cell proliferation such as the ventricular zone of the brain and the ependymal layer surrounding the central canal of the spinal cord. Also, although the mRNA is mostly in the neuroepithelium of the embryonic brain and spinal cord, the protein is widely distributed in these tissues, presumably as a consequence of transport in or along glial processes, local secretion, and/or redistribution as a consequence of cell migration (Engel et al., 1996; Meyer-Puttlitz et al., 1996). Based on the punctual expression pattern of phosphacan throughout the developing nervous system, it has been proposed that it may play a role in neuronal migration, differentiation, and circuit formation (Maeda and Noda, 1996; Meyer-Puttlitz et al., 1996).

Studies of DSD-1-PG localization in mouse brain match those for phosphacan in rat. For example, at E17 it is expressed in the dorsal, dorsolateral, and lateral subventricular zones and the cortical subplate (Gates et al., 1995); in the P7 cerebellum, it is strongly expressed in all layers, except the external granule layer, where it is only found on the Bergmann glia fibers (our unpublished observations) (Rauch et al., 1991). Preliminary studies of P6 cerebellar neurons plated on PORN and DSD-1-PG also suggest a neurite outgrowth promotion by the PG (our unpublished observations).

The effects on neurite outgrowth observed with hippocampal neurons and DRG explants in culture can also be correlated with the in vivo expression of DSD-1-PG/phosphacan. In E19 rat brain,
the strongest expression of phosphacan is in the marginal zone, subplate, and hippocampus (Meyer-Puttlitz et al., 1996). Furthermore, the DSD-1-epitope has been detected in the dentate gyrus of the rat hippocampus (Deller et al., 1997).

In the developing spinal cord, phosphacan is present in the growth territories of DRG axons. It is widely and strongly expressed by E13, being present in the dorsal root ganglion, the dorsal root entry zone, and the ventral roots, throughout the gray and white matter, and in the dorsal and ventral nerve roots present in the roof plate but not the floor plate (Meyer-Puttlitz et al., 1996). This pattern at E13 continues at E16 and E19 with more phosphacan in the dorsal, lateral, and ventral funiculi, in the motor columns, and the dorsal horn gray matter. On the basis of this distribution pattern, it has been suggested that phosphacan may be the PG that is associated with the inhibitory glial barriers to axonal extension at the dorsal root entry zones and in the roof plate of the spinal cord and that it may play a specific role in axonal growth and movement in the spinal cord by being in these glial-bordered extracellular spaces (Meyer-Puttlitz et al., 1996; Golding et al., 1997).

Similarly, several studies have reported the enhanced expression of growth-inhibiting CSPGs in the context of CNS lesions (McKeon et al., 1991; Bovolenta et al., 1993; Pindzola et al., 1993; Lips et al., 1995), and DSD-1-PG has also been found to be strongly upregulated after wounding in the CNS (Laywell et al., 1990; Laywell and Steindler, 1991; Barker et al., 1996; Deller et al., 1997), as has phosphacan (Snyder et al., 1996).

A study of the role of CSPGs in the outgrowth and adhesion of thalamic neurons plated onto living slices of the mouse embryonic neocortex (Emmerling and Lander, 1996) demonstrated that CS digestion could affect both the permissive environment of the subcortical plate and the neurite-repellent properties of the cortical plate. However, they suggest that the opposing activities of these different zones is attributable to differentially localized CS-binding factors rather than association with the presence of different CSPGs. As we show here, however, neurons from different regions of the CNS do not necessarily respond in the same way to the same CSPG. Hence, localized variations in the composition of the ECM may not always be required to obtain cell type-specific responses.

In the adult rat brain, it has been shown that DSD-1-PG occurs in perineuronal nets around parvalbumin-immunoreactive interneurons of the cerebral cortex because it was detected in the circumfluence of a selected subpopulation of neurons that expressed the calcium-binding protein parvalbumin occupying the extracellular space in close vicinity to the cell body and surrounding axon terminals and glial end feet but not the synaptic clefts (Wintergerst et al., 1996). It has been suggested that CSPGs associate with hyaluronic acid in such perineuronal nets or pericellular matrices to form a neuronal ECM structure analogous to that found in connective tissue (Maleski and Hockfield, 1997; Rauch, 1997). Different neuronal subsets have different complements of CSPGs (Celio and Blumcke, 1994) such that perineuronal CSPGs could regulate the extracellular milieu of neurons in cell type-specific ways. For example, late in development, the mature ECM may be an important element in limiting synaptic plasticity (Hockfield et al., 1990).

REFERENCES


