Nogo-A Inhibits Neurite Outgrowth and Cell Spreading with Three Discrete Regions

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Nogo-A is a potent neurite growth inhibitor in vitro and plays a role both in the restriction of axonal regeneration after injury and in structural plasticity in the CNS of higher vertebrates. The regions that mediate inhibition and the topology of the molecule in the plasma membrane have to be defined. Here we demonstrate the presence of three different active sites: (1) an N-terminal region involved in the inhibition of fibroblast spreading, (2) a stretch encoded by the Nogo-A-specific exon that restricts neurite outgrowth and cell spreading and induces growth cone collapse, and (3) a C-terminal region (Nogo-66) with growth cone collapsing function. We show that Nogo-A-specific active fragments bind to the cell surface of responsive cells and to rat brain cortical membranes, suggesting the existence of specific binding partners or receptors. Several antibodies against different epitopes on the Nogo-A-specific part of the protein as well as antiserum against the 66 aa loop in the C-terminus stain the cell surface of living cultured oligodendrocytes. Nogo-A is also labeled by nonmembrane-permeable biotin derivatives applied to living oligodendrocyte cultures. Immunofluorescent staining of intracellular, endoplasmic reticulum-associated Nogo-A in cells after selective permeabilization of the plasma membrane reveals that the epitopes of Nogo-A, shown to be accessible at the cell surface, are exposed to the cytoplasm. This suggests that Nogo-A could have a second membrane topology. The two proposed topological variants may have different intracellular as well as extracellular functions.

Key words: Nogo; reticulon; inhibitory regions; active sites; membrane topology; neurite outgrowth

Introduction

Regenerative nerve fiber growth and structural plasticity are limited in the adult mammalian CNS, in part because of the presence of neurite growth inhibitory constituents (Schwab and Bartholdi, 1996; Behar et al., 2000). An important step in elucidating the mechanisms mediating this inhibition was the molecular characterization of nogo-A, which encodes an oligodendrocyte-associated neurite growth inhibitor (Spillmann et al., 1998; Chen et al., 2000; GrandPré et al., 2000; Prinjha et al., 2000). The nogo gene gives rise to three major protein products, Nogo-A, -B, and -C, by both alternative splicing and alternative promoter usage (Chen et al., 2000; Oertle et al., 2003b). All Nogo isoforms share a common C-terminus of 188 amino acids, called reticulon-homology domain (RHD; Pfam PF02453) because of its similarity with the Reticulon (RTN) protein family (Roebroek et al., 1994, 1998; Moreira et al., 1999; Oertle et al., 2003c). Outside of this RTN domain, Nogo and the other three RTN genes have no obvious sequence similarities. Two long hydrophobic stretches (35 and 36 aa), which could serve as transmembrane (TM) domains and are probably responsible for the endoplasmic reticulum (ER) association of the proteins, are located in the RHD (van de Velde et al., 1994; Oertle et al., 2003a).

Myelin, oligodendrocytes, rat NI-250/NI-35 (for neurite outgrowth inhibitor of M, 250 and 35 kDa) as well as purified bovine Nogo-A-ortholog bNI-220 were shown to be inhibitory for fibroblast spreading and neurite outgrowth and to induce growth cone collapse of rat dorsal root ganglion (DRG) and chick retinal ganglion cell (RGC) neurons (Caroni and Schwab, 1988; Bandtlow et al., 1993; Rubin et al., 1995; Loschinger et al., 1997; Spillmann et al., 1998). The identification of the regions of Nogo-A that exert these diverse inhibitory effects in vitro and their possible accessi-
bility at the cell surface of oligodendrocytes would help us understand the mechanisms underlying the lack of regeneration in the CNS of higher vertebrates. Moreover, this information is crucial for the identification of Nogo-interacting molecules and for the development of optimal reagents for the neutralization of Nogo proteins. Potent neurite growth inhibitory activity was found in the Nogo-A-specific part of the molecule (Chen et al., 1999; Oertle et al., 2000; Prinjha et al., 2000). However, Nogo-A does not appear to be a conventional type I membrane protein; it does not possess an N-terminal hydrophobic sequence that could serve as a signal peptide such as is common to proteins that are secreted or expressed at the cell surface.

GrandPré et al. (2000) presented evidence that the 66 amino acid residue region (termed Nogo-66) between the two hydrophobic stretches of the RTN domain induces growth cone collapse and is exposed on the surface of Nogo-A-transfected cells. The cloning of a glycosyl phosphatidyl inositol (GPI)-anchored receptor (NgR) has been described recently that interacts with the Nogo-66 peptide (Fournier et al., 2001) as well as with the CNS myelin proteins OMgp (oligodendrocyte-myelin glycoprotein) and MAG (myelin-associated glycoprotein), both of which also have neurite growth inhibitory activity (Domeniconi et al., 2002; Liu et al., 2002; Wang et al., 2002a). The observation that Nogo-66 is inhibitory implies that all three Nogo isoforms should exert neurite growth inhibitory properties.

In the present study we provide evidence that three discrete regions of Nogo-A exhibit different inhibitory properties in vitro. Binding of Nogo-A-specific fragments to brain cortical membranes and the surface of responsive cells strongly argues for the existence of Nogo-A-specific receptor molecules. Antibody studies are suggestive for the expression of Nogo-A at the cell surface of cultured oligodendrocytes and of 3T3 fibroblasts and would imply that the N-terminal region of Nogo-A as well as the Nogo-66 region can face the extracellular space. A large intracellular pool of Nogo-A is associated with the endoplasmic reticulum (ER) and Golgi complex as revealed by double staining with marker proteins for these two organelles. Selective permeabilization studies suggest that a major part of the intracellular Nogo-A exhibits a second topology in which the N-terminus and Nogo-A-specific part of the molecule are exposed at the cytoplasmic side of membranes.

Materials and Methods


GST-Nogo-66 (aa 1055–1120 of human Nogo-A) has been cloned by PCR on human Nogo-A as a template.

Deletion constructs were cloned into pET28 vector (Novagen), pGEX-6P (Amersham Biosciences) and pET26 vector (Novagen). Human GST-Nogo-66 corresponds to the GST-nogo protein published by GrandPré et al. (2000). Synthetic rat peptide 4 EELVKQKYSNSALGHVN-STIKELRL corresponds to the human peptide 4 (ibid.) with one mismatch. Synthetic Pro/Ser-rich peptide (PSSPPSSPPSSPPSPSS) as well as rat peptide 4 have been produced and HPLC-purified by Primm SA. P472 (NYESIKHEPENPYPYEAA) was synthesized and purified by Research Genetics.

Production of recombinant Nogo-A-deletion library. The bacterial Nogo-A-deletion library was expressed in Escherichia coli. Proteins were extracted by repeated sonication in sonication buffer (20 ms Tris, 50 mM NaH2PO4, 100 mM NaCl, pH 8.0) with 0.75 mg/ml Lysozyme, by solubilization with B-Per (Pierce), or with 8 M urea. NiG expressed with pelb-leader was obtained from the periplasmic space according to the Nova-

Production of recombinant Nogo in Chinese hamster ovary cells. A 3119 bp fragment resulting from a partial HinclI digest of rat Nogo-A cDNA, NiR-G, was cloned into pSecTag2 expression vectors (Invitrogen, Groningen, The Netherlands). Transfection of pNiR-G into Chinese hamster ovary (CHO) cells resulted in intracellular, cytoplasmic expression of NiR-G. Stable NiR-G CHO cell lines were selected with 250 μg/ml Zeocin (Invitrogen). Recombinant NiR-G from cell lysate was purified over a Ni2+-NTA column (Qiagen AG, Basel, Switzerland).

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Cloning of rat NrG and stable Nr expressing CHO cell line. Adult rat brain poly-A + RNA was prepared using the Direct Quick Messenger RNA kit (Talent) according to manufacturer protocol. cDNA was prepared from 250 ng of poly-A + RNA with Moloney murine leukemia virus reverse transcriptase polymerase and poly-dT primers from Novagen. cDNA (1 μg) was used as a template for a PCR of 35 cycles with 5′-GTCCTGGATCCAGTAGAAGGGGCGTCC-3′ and 5′-GTTTCAGGTCAGGGGCGCCCGAGCAGCAGC-3′ as forward and reverse primers, respectively. The PCR product was subcloned into the BamHI-Xhol sites of pBluescriptII-KS and fully sequenced (Microsynth GmbH, Balgach, Switzerland).

pIg-NTAG-5-NgR was derived by subcloning rat NgR lacking the signal
Radioactive labeling and binding experiments. IMAC-purified NiG-Δ20 was iodinated by ANAWA Trading SA (Wangen, Switzerland) (2030 Ci/mmol) using Lactoperoxidase and purified by reverse-phase HPLC. Membranes from rat brain cortex were prepared as described (Olpe et al., 1990). Binding was performed for 1 hr at room temperature essentially as described (Kaupmann et al., 1997) using 1.5 ml tubes preincubated for 2 hr with 1% (w/v) bovine serum albumin to reduce nonspecific binding. Membrane homogenates in HEPES buffer, pH 7.4 (125 mM NaCl, 5 mM KCl, 0.6 mM MgCl₂, 1.8 mM CaCl₂, 20 mM HEPES, 6 mM dextrose), containing protease inhibitors (Roche Diagnostics, Mannheim, Germany) were incubated with 1.3 nM iodinated NiG-Δ20 in the absence or presence of increasing concentrations of unlabeled NiG-Δ20.

Antibodies. Rat and bovine antisera (AS) 472 were produced against the synthetic peptide NYESIKHEPENPPPYEA (bovine sequence) or the corresponding rat sequence SYDSIKLEPENPPPYEA (aa 623–640) (Chen et al., 2000) (Research Genetics Huntsville, AL). AS 922 was raised against the peptide RIYKGVQIAQKSDEGHPFRALEYEASEVIEMLVQKYSNASLGHY (aa 1055–1099 of human Nogo-A) from the Nogo-66 region, and AS 294 was raised against the C-terminal Nogo-A peptide KDAMAKIQAKIPGLKRKAD (Research Genetics). AS Rosa and AS Bianca were produced by intradermal immunization of prokaryotically produced, IMAC-purified, and gel electroeluted NiR-a and NiR-B. AS Bianca antibody (mAb) 11C7, whereas mAbs 11A8, 3D11, and 7B12 were produced against recombinant prokaryotically produced NiR-G. Spleen cells were fused with myeloma cells, and monoclonal lines were selected and subcloned. Supernatants of the obtained clones were screened on ELISA plates coated with the Nogo-A deletion library to localize their epitopes.

Rabbit anti-NgR antisera (AS α–NgR) were raised against three synthetic peptides of human NgR (EQQDSLATNQLRSVDPA, EVPCS-LPQLAGRDLKR, and GPRRPPGCSRKRNRTR) and affinity purified by Research Genetics.

Western blot analysis. SDS-PAGE and Western blotting were performed as described previously (Kelleher et al., 1992; Huber et al., 2002); blocking was done with 3% (w/v) Top Block (Juro Supply, Lucerne, Switzerland). Antibodies were diluted as follows: AS 472 1:2000; AS Bianca 1:10,000; AS Florina 1:20,000; affinity-purified AS 922 1.5 µg/ml; affinity-purified AS 294 0.3 µg/ml; affinity-purified AS α–NgR 0.11 µg/ml; α-BiP 2 µg/ml (Stressgen); mAb α-β-tubulin (Boehringer Mannheim, Mannheim, Germany) 1:200; mAb 9E10 α-myc (Invitrogen) 1:5000; monoclonal hybridoma culture supernatants 1:1500. Secondary antibodies were HRP-conjugated goat anti-rabbit (Pierce; 1:20,000) and anti-mouse (1:50,000).

Cell surface biotinylation. Brain oligodendrocyte cultures (van der Haar et al., 1998) were incubated with 2 mg EZ-LINK-Sulfo-NHS-LC-Biotin (0.25 mg/ml) (Pierce) per 75 cm² flask for 30 min at 20°C followed by incubation with 8 ml of 10 mM glycine in PBS with Ca²⁺/Mg²⁺ for 15 min to stop the biotinylation reaction and three washes with PBS. Cells were scraped, lysed in 1 ml lysis buffer [0.05 M NaH₂PO₄, pH 8.0, 0.15 M NaCl, 0.5% (w/v) CHAPS (Sigma), 2.5 mM iodoacetamide, 1 mM phenylmethylsulfonyl fluoride, 0.1 µg/ml aprotinin, 1 µg/ml leupeptin, 1 µg/ml pepstatin A] and precipitated three times with 100 µl of Dynabeads M-280 streptavidin (Dynal).

Immunocytochemistry. Optic nerve oligodendrocytes were prepared as described (Schwab and Caroni, 1988). Cultures (3–5 d old) grown on poly-l-lysine-coated coverslips were washed twice with PBS, fixed in 4% (w/v) paraformaldehyde (PFA), 5% (w/v) sucrose in PBS for 15 min at room temperature; cells were permeabilized with 0.1% (v/v) Triton X-100 (Tx-100) in PBS, and nonspecific binding was blocked with 10% (v/v) FCS. Cells were then incubated with affinity-purified AS 472 (1:200) or AS 922 (1:100) hybridoma culture supernatants containing mAb 11A8, mAb 3D11, and mAb 11C7 (all 1:100), anti-MG160 (1:100), and anti-calnexin (1:1000) for confocal microscopy in PBS, 1% FCS, 0.1% (v/v) Tx-100. Selective permeabilization of the plasma membrane was performed as described by De Strooper et al. (1997) with modifications. The cultures were washed twice with 10 mM PIPES buffer, pH 6.8, containing 0.3 M sucrose, 0.1 mM KCl, 2.5 mM MgCl₂, and 1 mM EDTA (De Strooper et al., 1997), followed by incubation of 30 min at room temperature in PIPES buffer with 12.5 µg/ml digitonin, containing the primary antibodies. Cells were washed with PIPES buffer and fixed and further processed. Secondary antibodies were goat anti-mouse tetrathymohydantoine isothiocyanate (TRITC) and goat anti-rabbit fluorescein-isothiocyanate (FITC) (Jackson ImmunoResearch Laboratories). A mouse monoclonal antibody against the KDEL sequence (α-BiP, 1:100; StressGen Biotechnologies, Victoria, British Columbia, Canada) was used to confirm the selective permeabilization of only the plasma membrane.

For cell surface staining, 2-d-old rat optic nerve cultures were incubated with AS 472 (1:500), AS 922 (1:500), AS Bianca (1:500), or monoclonal antibodies (undiluted supernatant) in medium for 25 min at room temperature. For CHO/oligodendrocyte co-cultures, 6000 CHO cells per well were added 24 hr before staining procedures. Cultures were washed, fixed with 4% (w/v) PFA and 5% (w/v) sucrose and blocked with 0.1 M NaCl, 0.25% (w/v) bovine serum albumin and 0.05% (v/v) Triton X-100 for 1 hr.
maleic acid with 2% (w/v) blocking reagent (Roche) for 1 hr. Secondary alkaline phosphatase-conjugated antibodies (Milan Analytica, Lausanne, Switzerland) were used at 1:7500 in 0.1 M maleic acid with 1% (w/v) blocking reagent (1 hr). The cultures were washed twice with maleic acid buffer and once with alkaline phosphatase buffer (0.1 M Tris-HCl, pH 9.5, 0.1 M NaCl, 5 mM MgCl₂), and the staining was developed for 3 hr at room temperature with 0.175 mg/ml 5-bromo-4-chloro-3-indolyl phosphate (Sigma) and 0.338 mg/ml nitroblue tetrazolium (Sigma) in alkaline phosphatase buffer. For surface MAG and intracellular tetrazolium (Sigma) in alkaline phosphatase buffer, the staining was developed for 2 hr instead of 3 hr.

The plgκ−V5-NgR CHO cells were washed, fixed for 15 min at room temperature, and blocked with 10% (v/v) FCS in PBS for 20 min at room temperature. mAb α-V5 (Invitrogen R-960–25) was diluted 1:300 in 1% (v/v) FCS in PBS for 1 hr and washed, and secondary FITC-conjugated anti-mouse antibodies (Jackson ImmunoResearch Laboratories) were used at 1:200 in PBS for 30 min.

Flow cytometry. Flow cytometry and cell sorting were performed on a Cytomation MoFlo high-speed cell sorter (Fort Collins, CO). The flow cytometer was equipped with an argon/UVP Enterprise II laser tuned to 488 nm with 130 mW of power. Fluorescein (FITC) fluorescence was collected through a 530/40 nm band-pass filter. For analysis, 3T3 fibroblasts were detached with Cell Dissociation Buffer (Invitrogen) or 0.5% (w/v) trypsin in PBS. The preformed complex used to detect binding of NiR-G to 3T3 fibroblasts was prepared as follows: NiR-G and anti-Myc antibody (9E10; Sigma) were incubated at a 1:1 molar ratio for 30 min at 4°C. Next, FITC-conjugated F(ab)₂, goat anti-mouse IgG was added and incubated for an additional 30 min at 4°C. The resulting molar ratio of the trimeric complex was 1:1:0.5. The complex was added to 10⁶ 3T3 fibroblasts in a final volume of 100 μl, incubated for 2 hr at 4°C, washed, and analyzed by flow cytometry.

In vitro assays. 3T3 fibroblast spreading assays were performed as described previously (Spillmann et al., 1998). CHO spreading assays were performed essentially the same way as for 3T3 fibroblasts. Briefly, CHO cells were split 1:2. Twenty-four hours later they were trypsinized in PBS-EDTA for 30 sec, and ~8000 CHO cells were plated onto culture dishes precoated with 5, 1, 0.5, and 0.2 μg per well NIG or Nogo-66. After 30–45 min, the cells were fixed with 4% (w/v) PFA, 5% (w/v) sucrose and then analyzed.

PC12 neurite outgrowth assays were performed as described previously (Rubin et al., 1995).

Neurite outgrowth assays with P7 rat cerebellar granule cells were performed as described by Niedero¨schinger, 1997; Fritsche et al., 1999).

Retinal ganglion cell stripe assays were performed according to Vielmetter et al. (1990) with modifications (Schmalfeldt et al., 2000). Exemplars were evaluated after fixation with 4% (w/v) PFA, 0.1% (v/v) glutaraldehyde in PBS for 10 min at room temperature. For immunostainings, fixed explants were blocked for 1 hr at room temperature with RNO-blocking solution [0.5% (w/v) BSA, 0.3% (w/v) Top-Blot (Juro Supply), 0.1% (w/v) NaN₃ in PBS], permeabilized for 10 min with 0.05% (v/v) Tx-100 in RNO-blocking solution, frozen for 1 min at −20°C, and incubated with primary antibodies (AS Bianca for NiR, AS Laura for Nogo-A, NiR-G, NIG, NIGΔ3, and NIGΔ20, and Novagen mAb anti-T7 for Nogo-C and β-Gal control protein). After washing with PBS, FITC- and TRITC-conjugated antibodies (Jackson ImmunoResearch Laboratories) were added (1:150) to the explants. The samples were coveredslipped in 50% (v/v) glycerol, 25 mM NaHCO₃, 40 mM NaCl, 1% (w/v) p-phenylenediamine (Sigma).

Growth cone collapse assays on chick and rat DRG neurons were performed essentially as described previously (Bandtlow et al., 1993; Bandtlow and Loschinger, 1997; Fritsche et al., 1999).

Figure 2. Analysis of the inhibitory properties of various Nogo fragments on 3T3 fibroblast spreading. The main inhibitory activity resides in two regions of the Nogo-A protein. Nogo-A deletion products were tested for their relative inhibitory activity (percentage inhibition) on fibroblast spreading (5 μg/100 μl, 1 μg/100 μl, 0.5 μg/100 μl, and 0.2 μg/100 μl of fragments coated per square centimeter). Mean of spreading on plastic is indicated by the dotted line. For each fragment at least three independent assays with proteins from at least two separate purifications were performed. The fragments that are most inhibitory for fibroblast spreading are highlighted. Control peptides (5 μg/100 μl coated per square centimeter) did not exhibit inhibitory properties except for l-laminin.
Results

The three main isoforms of Nogo, the most important domains and active sites, and the epitopes of the antibodies and antisera used in this study are summarized in Figure 1.

Two regions in the N-terminal part of Nogo-A are inhibitory for spreading of 3T3 fibroblasts

To identify the regions of Nogo-A responsible for the inhibition of 3T3 fibroblast spreading, a library of 50 Nogo deletion constructs was made, and recombinant proteins were expressed in bacteria (Fig. 2). The Co\textsuperscript{2+}-affinity chromatography-purified Nogo fragments were coated on tissue-culture dishes. The purified material of some of the recombinant proteins contained lower molecular weight Nogo fragments (Chen et al., 1999), because the Nogo-A cDNA sequence contains internal ribosomal binding sites for E. coli and numerous rare codons (our unpublished observations). The purity of fragments was estimated on the basis of silver-stained SDS-PAGE (see Fig. 6D). The comparison between the different deletion constructs is therefore semiquantitative.

The apparent EC\textsubscript{50} for inhibition of 3T3 fibroblast spreading was \(\sim 400-500\ ng/100\ \mu l\) Nogo-A coated overnight per cm\textsuperscript{2} of culture dish (\(\sim 4\ \text{pmol/cm}^2\)). Treatment of Nogo-A or its fragments with 8 M urea resulted in a strong reduction of inhibitory activity, indicating that conformation is important.

The analysis of Nogo fragments in the fibroblast spreading assay revealed that at least two stretches of the Nogo-A protein mediate inhibition of the spreading of freshly plated fibroblasts, namely aa 59–172 (NiR-\Delta\text{2}) and aa 544–725 (NiG-\Delta\text{20}) (Figs. 2, 3). All of the fragments derived from the Nogo-A-specific region (NiG) displaying inhibitory activity (e.g., NiG-\Delta\text{4} and NiG-\Delta\text{8}) partially overlap with aa 544–725 (NiG-\Delta\text{20}). Minor inhibitory activity at high protein concentration was seen for the C-terminal part of the Nogo-A-specific region (aa 763–865; NiG-\Delta\text{19}). At high concentrations, the anti-spreading activity of the Nogo fragments became anti-adhesive. These two effects could be overcome eventually by the cells after longer incubation times, indicating that they were not caused by toxic effects (data not shown). Nogo-C, Nogo-66, and Nogo-66 Peptide 4 (shown to be the inhibitory region of Nogo-66 by GrandPré et al. (2000)) were not inhibitory for fibroblast spreading (Fig. 2).

We found no correlation between the isoelectric point of the protein fragments and their inhibitory character. The high occurrence of proline and serine in the N-terminus of Nogo-A is also accompanied by strong axonal fasciculation. Control \(\beta\)-galactosidase/laminin stripes (stained with anti-neurofilament mAb) are permissive substrates for RGC axons. B, Inhibition score for RGC outgrowth on different Nogo fragments. Cultures were evaluated by giving a score of 5 for striped neurite outgrowth with no fibers crossing the Nogo containing stripes, 4–2 for striped neurite outgrowth with increasing numbers of crossing fibers, 1 for random neurite outgrowth with tendency to grow in the direction of the stripes, and 0 for complete random neurite outgrowth. The dotted line indicates the mean of all \(\beta\)-galactosidase/laminin stripes (stained with AS Laura; arrowhead). This avoidance is concentration dependent and accompanied by strong axonal fasciculation. Control \(\beta\)-galactosidase/laminin stripes (stained with \(\alpha\)-17 mAb) are permissive substrates for RGC axons. B, Inhibition score for RGC outgrowth on different Nogo fragments. Cultures were evaluated by giving a score of 5 for striped neurite outgrowth with no fibers crossing the Nogo containing stripes, 4–2 for striped neurite outgrowth with increasing numbers of crossing fibers, 1 for random neurite outgrowth with tendency to grow in the direction of the stripes, and 0 for complete random neurite outgrowth. The dotted line indicates the mean of all \(\beta\)-galactosidase/laminin stripes (stained with AS Laura; arrowhead). This avoidance is concentration dependent and accompanied by strong axonal fasciculation.
not responsible for the observed effect because a poly-Pro/Ser peptide is not inhibitory (Fig. 2, Control peptides).

These data show that the anti-spreading activity of Nogo-A on 3T3 fibroblasts resides in two defined stretches located at the N-terminus (aa 59–172) and within the Nogo-A-specific part (aa 544–725; NiG-H25) of the protein. Nonspecific physicochemical properties (acidity of the fragments, structural effects attributable to proline and serine residues) are not responsible for this effect. The C-terminal RTN-homology domain including Nogo-66 is not involved in the inhibition of fibroblast spreading.

Several regions of Nogo-A are inhibitory for neurite outgrowth

To determine whether the fragments of Nogo-A that were non-permissive for cell spreading are also inhibitory for neurite outgrowth, we tested a series of bacterially produced Nogo-A fragments as well as eukaryotically produced Nogo-fragment–alkaline phosphatase (Nogo-AP) fusion proteins in different neuronal assays.

In a substrate stripe assay (Vielmetter et al., 1990), we tested whether Nogo stripes are inhibitory for the growing embryonic chicken [embryonic day (E) 7–9] RGC axons. Neurites avoided laminin/Nogo-A-coated stripes, growing on the laminin-only stripes (Fig. 3A), whereas stripes coated with laminin/β3-galactosidase were not circumvented. Full-length Nogo-A was strongly nonpermissive for RGC neurites, whereas the N-terminal part of Nogo-A (-B (aa 1–172) had only marginal effects. Nogo-C activity was indistinguishable from the control protein β-galactosidase (Fig. 2A, B). The Nogo-A-specific region NiG-D20 (aa 544–725) appears to contain the main inhibitory region for these neurites (Fig. 3B). The growth cones were also seen to stop when entering NiG-D20-coated “dead-end” lanes (data not shown). These nonpermissive effects were concentration dependent: increasing numbers of crossing fibers were observed at lower concentrations of Nogo-A or its active fragments (Fig. 3A, B). No obvious difference was detected between nasal and temporal RGC neurites concerning their responsiveness to Nogo-A regions.

A laminin-independent, NGF-responsive clone of PC12 cells (Rubin et al., 1995) was primed with 50 ng/ml NGF for 24 hr and then plated onto dishes coated with bacterially produced Nogo fragments at 0.1–3 μg/cm². Neurite outgrowth was scored 1 d later. The Nogo-A-specific region (NiG) and its aa 544–725 fragment (NiG-D20) strongly inhibited PC12 neurite outgrowth (Fig. 3C, D). In contrast, the N-terminal fragment (aa 1–172) had only minor activity, detectable only at high protein concentrations. Nogo-C and Nogo-66 were inactive.

Substrate-coated NiG (aa 174–979) also acted as a strong neurite growth inhibitor of primary mammalian neurons: neurite outgrowth of postnatal day (P) 7 rat cerebellar granule cells was prevented in a dose-dependent manner, and cells aggregated at higher Nogo substrate concentrations (Fig. 3E).

We further tested the growth cone collapsing activities of the various Nogo constructs. Murine GST-Nogo-66, human GST-Nogo-66, and HPLC-purified Nogo-66 lacking the GST tag induced growth cone collapse of E13–E15 chicken DRG explants as described previously (GrandPré et al., 2000; Fournier et al., 2001). The growth cone collapsing activities of these proteins were relatively low (~50% collapsed at ~1 μM) and varied between preparations, probably because of the poor solubility of these recombinant proteins. Nogo-66 peptide 4 was inactive. In contrast, soluble dimeric Nogo-66-AP was collapse-inducing in a range similar to what has been described previously with an apparent EC50 of ~2 nM (Fig. 4B) (GrandPré et al., 2000). Growth cone collapse could also be elicited by the Nogo-A-specific, soluble, and dimeric fragment aa 544–725 (NiG-D20-AP), with an apparent EC50 of 200–400 nM (Fig. 4B). Bacterially expressed recombinant monomeric NiG (aa 174–979) or NiG-D20 (aa 544–725) had no collapse-inducing activities, nor had NiR-G (aa 1–979). Time-lapse studies on dissociated rat P6 DRG neurons revealed that growth cone collapse induced by Nogo-66-AP and NiG-D20-AP occurred with a slower time course than collapse elicited by hSema3A (Fig. 4A, C). In addition, although AP-Sema3A induced collapse of virtually all growth cones at low nanomolar concentration, both Nogo-66-AP and NiG-D20-AP did not lead to collapse of >60% of the rat or chicken growth cones (Fig. 4A, B).
The Nogo-66 receptor NgR is not required for inhibition of cell spreading

We isolated a cDNA encoding the rat NgR protein (GenBank accession number AF462390) and generated antisera against specific peptides of the protein. To study the role of NgR for inhibition of cell spreading, we generated a CHO cell line stably expressing rat NgR at high levels on its cell surface (Fig. 5A, B). Wild-type CHO cells (CHO-wt) do not express detectable amounts of NgR as determined by RT-PCR (40 cycles; data not shown) and Western blotting (Fig. 5B). Spreading of both CHO-wt and CHO-NgR was dose-dependently inhibited by Nogo-A aa 174–979 (NiG). Independent of NgR expression, neither cell line responded to coated Nogo-66 (Fig. 5C).

These results show that the inhibition of fibroblast-like cells by the Nogo-A-specific active region can occur in the absence of NgR and that the presence of NgR does not change the responsiveness of the cells to Nogo-66 or Nogo-A.

Presence of binding site(s) for active Nogo-A-specific fragments on 3T3 fibroblasts and rat brain cortical membranes

Because the Nogo-A fragments aa 1–172 and aa 544–725 (NiR-Δ2 and NiG-Δ20) were inhibitory for cell spreading and neurite outgrowth despite the absence of NgR and independently of NgR, the presence of a separate, Nogo-A-specific receptor has to be postulated. We therefore performed binding studies of multimerized, myc-tagged, and IMAC-purified Nogo-A aa

Figure 5. Activity of NgR-expressing CHO cells versus wild-type cells on Nogo fragments. A, An mAb against the V5-epitope stains the surface of a CHO cell line stably transfected with rat NgR. B, On Western blot, no NgR protein is detectable in lysates of wild-type CHO cells, whereas high amounts of NgR are detectable in lysates of the CHO–NgR cell line. C, Both wild-type and NgR-transfected CHO cells are strongly inhibited in spreading by the Nogo-A-specific region NiG, whereas they are unresponsive to Nogo-66.

Figure 6. Binding of amino-terminal Nogo fragments to 3T3 cells and cortical membranes. A, Binding of NiR-G (aa 1–979) to 3T3 cells. 3T3 fibroblasts were incubated with a preformed complex consisting of α-myc antibody (9E10), FITC-conjugated F(ab)_2 goat α-mouse IgG fragments, and myc-tagged amino-terminal fragment of Nogo-A (NiR-G). Binding of the complex to 3T3 fibroblasts was analyzed by flow cytometry. Unstained 3T3 fibroblasts were used as negative control. Further negative controls include 3T3 cells incubated with a complex consisting of the α-myc antibody and the FITC-conjugated α-mouse Ab or with the FITC-conjugated α-mouse Ab alone. B, Binding of NiR-G to 3T3 cells is protease sensitive: trypsinization of 3T3 fibroblasts before incubation with Nogo completely abolishes Nogo-binding to their surface. C, Western blot of purified, myc-epitope-tagged NiR-G. D, Silver-stained gel of IMAC-purified NiG-Δ20. M, Molecular weight standard. E, Binding of 1.3 nm [%125I]-NiG-Δ20 to rat brain cortical membranes and competition by increasing concentration of unlabeled NiG-Δ20. □ indicates the values obtained after incubation of 1.3 nm [%125I]-NiG-Δ20 in the absence of cortical membranes, i.e., nonspecific binding to the tubes. Values are represented as mean ± SE (**p < 0.01; ***p < 0.001; Student’s t test for competition; n = 3).
1–979 (Fig. 6C) to living 3T3 fibroblasts. The cells were labeled in suspension at 4°C and subsequently analyzed by flow cytometry. Anti-myc monoclonal antibody-complexed myc-tagged Nogo-A fragment (NiR-G) bound efficiently to 3T3 cells as seen by a shift in fluorescence of >90% of the 3T3 cells (Fig. 6A). In contrast, 3T3 cells were not labeled after incubation with the primary mouse anti-myc mAb complexed with an FITC-conjugated secondary F(ab)2 goat anti-mouse IgG in the absence of Nogo-A or with the secondary Ab alone. The binding of the amino terminal Nogo-A fragment to the surface of fibroblasts is protease sensitive, because 3T3 cells that were dissociated with trypsin before Nogo-A incubation do not bind Nogo-A (Fig. 6B).

To test binding of Nogo-A aa 544–725 (NiG-Δ20) to rat cortical membranes, we used [125I]-labeled NiG-Δ20 in a radioligand binding assay (Fig. 6D). [125I]-labeled Nogo fragment bound to these brain plasma membranes. The specificity of binding was shown by a concentration-dependent competition of [125I]-NiG-Δ20 binding at a concentration of 1.5 nM by increasing amounts of unlabeled Nogo-A aa 544–725 (Fig. 6E).

These results show that defined, bioactive fragments of Nogo-A can bind to the surface of 3T3 cells and rat cortical membranes, demonstrating the presence of membrane-bound, Nogo-A-specific binding sites or receptor(s) different from the Nogo-66 receptor NgR.

Antibodies against Nogo-A
Antisera and monoclonal antibodies were raised against Nogo-A peptides or recombinant protein fragments (Fig. 1). The monoclonal antibody 11C7 was raised against Nogo-A aa 623–640, i.e., the same Nogo-A specific peptide as the rabbit AS 472 (Chen et al., 2000). Three other mAbs raised against Nogo-A aa 1–979 (NiR-G) reacted with different epitopes as determined by ELISA on the Nogo protein fragment library: 11A8 recognizes the most N-terminal part of the Nogo-A-specific region (aa 209–233), and 7B12 (aa 763–820) and 3D11 (aa 910–920) recognize the C-terminal part of the Nogo-A-specific region of the molecule. Two antisera (AS Bianca and AS Rosa) recognize the N-terminal region aa 1–172 (NiR), common to both Nogo-A and Nogo-B. Finally, antisera were raised against the Nogo-66 region (AS 922) and the C-terminus of Nogo including the ER retention signal (AS 294), respectively.

Antibodies against all epitopes identified the 190 kDa Nogo-A band on a Western blot of oligodendrocyte cell culture homogenate (Fig. 1B). AS Bianca, AS 922, and AS 294 also recognize the 55 kDa band of Nogo-B. mAb 3D11 and AS 922 recognized an additional band at ~80 kDa, which could be a breakdown product of Nogo-A. At longer exposure times, AS 922 and AS 294 both stained additional bands even after affinity purification.

Figure 7. Nogo-A is present at the cell surface of cultured oligodendrocytes. Live, unpermeabilized oligodendrocytes (3 d in culture) were incubated with anti-Nogo-A antibodies, fixed, and visualized with secondary antibodies conjugated with alkaline phosphatase. All antibodies against the N-terminus of Nogo-A stain the cell surface of oligodendrocytes (mAb 11C7, mAb 11A8, mAb 3D11, mAb 7B12, AS Bianca), although more weakly than an antibody against the surface glycoprotein MAG (shorter development; see Materials and Methods). The control mouse IgG (d) and an antibody against the intracellular oligodendrocyte protein CNPase (e) do not stain oligodendrocytes. Preincubation of 11C7 with the immunogen, peptide P472, reduced the staining to background levels (b), whereas preincubation with an unspecific peptide (Px) did not result in a reduction in staining intensity (c). Furthermore, AS 922 against the C-terminal Nogo-66 region stains the cell surface of living oligodendrocytes (f), whereas the corresponding preimmune serum (pre-922) gives only background staining (m). Staining of co-cultures of oligodendrocytes with CHO cells expressing rat Nogo-A with mAb 11C7 shows that although intact oligodendrocytes are stained, Nogo-A produced by CHO cells (arrow) is not detectable at the cell surface (g). Both cell types are stained with 11C7 after permeabilization (p). Scale bar, 25 μm.

Domains of Nogo-A present at the cell surface of cultured oligodendrocytes
Cultures of unpermeabilized living oligodendrocytes were incubated with different anti-Nogo-A antibodies. The Nogo-A-specific mAbs 11C7, 11A8, 7B12, and 3D11 (Figs. 1, 7a,f,g,h) as well as AS 472 (data not shown) stained the surface of the differentiated oligodendrocyte cell bodies and their process network. Compared with stainings for GalC or sulfatide (data not shown) or an mAb against the oligodendrocyte surface protein MAG (Fig. 7n), the Nogo staining was relatively weak. The specificity of the cell surface staining is suggested by the following results. (1) The control mouse IgG and the antibodies against the intracellular protein CNPase did not stain intact living cells (Fig. 7d,e). (2) Preincubation of mAb 11C7 (or AS 472; data not shown) with the corresponding immunogenic peptide P472 reduced staining to background levels (Fig. 7b). Preincubation of mAb 11C7 or AS 472 with an unspecific peptide (Px) before their addition to the cells did not reduce the staining (Fig. 7c). (3) Co-cultures of oligodendrocytes with transfected Nogo-A-expressing CHO cells were stained with mAb 11C7. Although intracellular Nogo-A was detectable in both cell types after permeabilization (Fig. 7p), surface Nogo-A could be detected only on oligodendrocytes but not on the transfected CHO cells (Fig. 7o). This observation suggests that transport of Nogo-A to the cell surface could be cell-type
specific. AS Bianca recognizing the N-terminus of Nogo-A, but not the corresponding preimmune serum, also stained the cell surface of living oligodendrocytes (Fig. 7i,k). Staining with AS 922 showed that also the Nogo-66 region is exposed on the surface of oligodendrocytes, confirming previous reports (GrandPre et al., 2000). Because AS 922 stains additional bands on Western blot (Fig. 1B), however, we cannot exclude the possibility that the antiserum also recognizes proteins other than Nogo on the cell surface of oligodendrocytes. Cell surface staining was present on all major and small processes and on the cell body.

These results suggest that the Nogo-A-specific part and the N-terminus of the molecule as well as the Nogo-66 loop are exposed to the extracellular space on the plasma membrane of oligodendrocytes.

Cell surface biotinylation of Nogo-A

To confirm the presence of Nogo-A on the cell surface of oligodendrocytes, living, unpermeabilized cultures of newborn rat brain oligodendrocytes were exposed to a membrane-impermeable biotinylating reagent. Biotinylated proteins were precipitated with streptavidin-coated beads. The precipitated proteins and the supernatants of the precipitates were analyzed by Western blot. In the precipitate, Nogo-A was recognized by the Nogo-A-specific AS 472 with the expected molecular weight of 190 kDa (Fig. 8). This band was also recognized by another anti-Nogo-A antiserum, AS Bruna (data not shown) and was not detected when biotin was omitted in control experiments (data not shown). The abundant intracellular proteins β-tubulin and BiP were not present in the precipitated material but were detected in the supernatant (Fig. 8). In line with the data shown below (see Fig. 10), large amounts of Nogo-A were also present intracellularly. Densitometric analysis of blots of three separate experiments revealed that ~1% of total cellular Nogo-A could be precipitated by biotinylation from the cell surface of the cultured oligodendrocytes.
Nogo-A is present at the surface of 3T3 fibroblasts with Nogo-A-specific regions facing the extracellular space

Nogo-A is expressed endogenously in various cultured cell lines including the Nogo-A-responsive 3T3 fibroblasts (as shown by RT-PCR and Western blot) (Oertle et al., 2003b). To analyze whether Nogo-A-specific regions are also present on the surface of these cells, 3T3 fibroblasts were detached from the culture flask with Cell Dissociation Solution and incubated in suspension with 0.1 μM Nogo-A-specific mAb 11C7, followed by FITC-conjugated secondary Ab. Cells that bound the Nogo-A antibody were separated from unlabeled cells by fluorescence-activated cell sorting. mAb 11C7, but not a mouse IgG control Ab or the secondary Ab alone, labeled the 3T3 cells as seen by a large shift of the fluorescence intensity of the 3T3 cell population (Fig. 9).

This result indicates that the mAb 11C7 epitope, aa 623–640, which lies within the Nogo-A-specific region with highest inhibitory activity, is present on the surface of living 3T3 fibroblasts.

Intracellular localization and topology of Nogo-A

Fixed and permeabilized oligodendrocytes in cultures derived from P7–P10 optic nerve were stained with AS 472, AS 922, AS Bianca, and the mAbs 11C7, 11A8, and 3D11 (Fig. 10). The bulk of the staining was reticular, present in both the cell body and the large processes of the oligodendrocytes. Note the difference in staining between Nogo-A (Fig. 10, 11C7 Tx-100), an ER membrane protein, and BiP (Fig. 10, α-Bip Tx-100), which is a soluble protein present in the ER lumen. (The same reticular staining patterns were obtained with calnexin, also an ER membrane protein) (Fig. 11 A–C). Nogo-A was not detectable on the membrane sheaths formed between the fine processes at longer culture times (data not shown). To elucidate the orientation of the intracellular protein, we selectively permeabilized oligodendrocytes with digitonin (DIG), an agent that at low concentrations permeabilizes solely the plasma membrane and leaves intracellular membranes intact (De Strooper et al., 1997). In cultures permeabilized with DIG, all antibodies and antisera except for AS 922 stained Nogo-A in the cytoplasm of oligodendrocytes (Fig. 10, right row). Staining with an antibody against a lumenal ER protein, BiP, was negative (Fig. 10), whereas staining against actin was positive (Fig. 10), showing the selectivity of the plasma membrane, but not ER, permeabilization.

These results demonstrate that the Nogo-A-specific region and the N-terminus of Nogo-A are exposed to the cytoplasmic side of intracellular membranes, at least for a large part of the Nogo-A residing in the ER.

We then examined in which intracellular compartment Nogo-A was present by colocalization studies using confocal microscopy. Double labeling with antibodies against Nogo-A and the ER marker calnexin (Fig. 11 A–C) revealed that most but not all Nogo-A colocalized with calnexin. The colocalization was most marked in the oligodendrocyte processes. In the cell body, an area could often be observed to one side of the nucleus where only Nogo-A was present but the ER marker was absent (Fig. 11C, arrowhead). In this perinuclear region, Nogo-A colocalized with the Golgi marker MG160 (Gonatas et al., 1989; Croul et al., 1990; Gonatas et al., 1995) (Fig. 11 D–F). Thus, Nogo-A is present intracellularly in both the ER and the Golgi complex.
Discussion

Three domains of Nogo-A are differentially involved in the inhibition of cell spreading, neurite outgrowth, and growth cone collapse

More than 50 Nogo protein fragments were expressed in E. coli or eukaryotic cells and coated as culture substrates or added as soluble proteins to cultured cells. Three active regions of Nogo could be defined. (1) On the basis of our semiquantitative analysis, the N-terminus of Nogo-A--B inhibits 3T3 cell spreading as a substrate but has only a minor effect on neurons. (2) The Nogo-A-specific region and its central 181 amino acids are an inhibitory substrate for primary neurons, PC12 cells, and fibroblasts, and the same regions induce growth cone collapse when added as dimers in solution. (3) Dimeric Nogo-66 (loop region between the two C-terminal hydrophobic regions) induces growth cone collapse, but Nogo-66 or Nogo-C coated as substrate are inhibitory for neither E7--E9 chick retina or PC12 cell neurite outgrowth nor fibroblast spreading. Thus, Nogo-66 peptide seems to be prime facie more potent than the Nogo-A-specific region in inducing DRG growth cone collapse, whereas the Nogo-A-specific region has a stronger neurite outgrowth inhibitory activity. This might indicate that the interaction of Nogo-66 with its receptor NgR could be involved primarily in axonal guidance, whereas the Nogo-A-specific region could be involved in limiting plasticity and regeneration. It is known from other proteins that multiple distinct regions might contribute to their inhibitory activity (Ugrhin et al., 2003).

The potent inhibitory activity of the Nogo-A regions has been recognized previously (Chen et al., 1999, 2000; Oertle et al., 2000; Prinjha et al., 2000; Fournier et al., 2001). The physiological importance of this region is emphasized by the fact that an antiserum (AS 472) against aa 623--640 in the Nogo-A-specific region neutralizes the inhibitory activity of CNS myelin in vitro (Chen et al., 2000) and induces sprouting of adult rat Purkinje axons in vivo (Buffo et al., 2000). A contribution of the aa 763--865 region (NiG-D19) to the inhibitory activity in vivo is emphasized by recent findings showing that pump-infused mAb 7B12, which binds to an epitope within NiG-D19, leads to compensatory axonal sprouting along with improved functional recovery after experimental stroke in adult rats (Wiessner et al., 2003). On the other hand, the role of Nogo-B and Nogo-C (both containing Nogo-66), which have a broad expression pattern in peripheral tissues as shown by Northern and Western blots (Morris et al., 1999; Tagami et al., 2000; Huber et al., 2002), is still unclear.

The Nogo-A-specific region has an inhibitory effect on a range of cells including neurons, 3T3 fibroblasts, and CHO cells. Nogo-A, which is predominantly present in oligodendrocytes and myelin in the adult CNS, may therefore not only restrict neurite growth, plasticity, and axonal regeneration, but also limit the invasion and migration of cells and tumors in the CNS (Amberger et al., 1998; Belien et al., 1999). The presence of Nogo-A in various cell lines in vitro (Oertle et al., 2003b), and on the surface of 3T3 fibroblast cells in particular, may suggest an additional role, e.g., in the context of cell contact-mediated growth control.

All of these results suggest that the Nogo-A molecule possesses several binding sites (at least two for fibroblasts and neurons, respectively) and that the Nogo receptor is possibly a complex composed of several different subunits. One such subunit, NgR, serves as receptor component for the Nogo-66 peptide (Fournier et al., 2001), OMgp (Wang et al., 2002a), and MAG (Domeniconi et al., 2002; Liu et al., 2002). Being a GPI-anchored protein, it requires additional subunits for signal transduction. The bioactive Nogo-A-specific regions bind efficiently to the surface of 3T3 fibroblasts and to rat brain membranes, hence strongly suggesting the presence of Nogo-A-specific receptor(s). Our data demonstrate that the Nogo-A-mediated inhibition of cell spreading can occur in the absence of NgR and that the presence of NgR does not change the responsiveness of CHO cells to Nogo-66 or Nogo-A.

Activation of the small GTPase RhoA has been shown to be a crucial step in the signal transduction of inhibitory cues in various neurons (Li et al., 2002; Winton et al., 2002). The Nogo-66 peptide can activate Rho-A (Niederost et al., 2002), probably via activation of the NgR coreceptor p75NTR (Wang et al., 2002b). Interestingly, the Nogo-A fragments aa 174--979 (NiG) and aa 544--725 (NiG-D20) have been shown to also activate RhoA and inhibit Rac in cerebellar granule cells and 3T3 fibroblasts (Niederost et al., 2002), and this still occurs after removal of NgR by phosphoinositide-specific phospholipase C treatment. These results suggest that RhoA activation is a key downstream component for both NgR and the putative Nogo-A-specific receptor.

Heteromeric receptor complexes interacting with different binding sites of a given ligand are well known for axonal guidance molecules [e.g., plexins and neuropilins for semaphorins (Tama-
gnone et al., 1999]) as well as for neurotrophic factors [e.g., tropomyosin related kinases and p75 neurotrophin receptor (p75NTR) for neurotrophins (Lee et al., 2001)] and inhibitory proteins [e.g., p75, NgR and gangliosides for MAG (Domeniconi et al., 2002; Liu et al., 2002; Vyas et al., 2002; Yamashita et al., 2002)]. Whether the specificity of the response to Nogo (growth inhibition vs growth cone collapse; cell type specificities) is linked to specific binding sites and receptor subunits remains to be investigated.

Inhibitory regions of Nogo-A are exposed at the cell surface of oligodendrocytes

Several lines of evidence suggest the presence of large parts of Nogo-A, including the inhibitory regions at the cell surface facing the extracellular space. These include cell surface biotinylation, immunocytochemistry on living cultured oligodendrocytes with three antisera and four monoclonal antibodies, all against different parts of Nogo-A, and cell sorting of surface-labeled 3T3 fibroblasts. The presence of Nogo-A at the cell surface of oligodendrocytes in vitro is consistent with results obtained in vivo: sprouting of axons can be elicited in intact regions of the adult rat CNS by application of AS 472 or mAb 7B12 (Buffo et al., 2000; Wiessner et al., 2003).

Nevertheless, a high proportion of Nogo-A protein was found intracellularly, associated with ER and Golgi membranes. In transfected cell lines such as COS or CHO cells, ectopically expressed Nogo is also localized in the ER and Golgi and was not detected on the cell surface (data not shown). Interestingly, Nogo-A was detected on the cell surface only in differentiated oligodendrocytes, but intracellular Nogo-A was present also in oligodendrocyte precursors (Wang et al., 2002c) (our unpublished observations). This is in agreement with the observation that the contact-mediated inhibition exerted by cultured oligodendrocytes was not detectable in precursor cells (Schwab and Caroni, 1988; Bandtlow et al., 1990). In addition, intracellular Nogo-A released to the extracellular matrix after oligodendrocyte and myelin damage might contribute to the inhibition of axonal regeneration as has been proposed previously (Goldberg and Barres, 2000).

Permeabilization of only the plasma membrane of oligodendrocytes before addition of antibodies revealed that the Nogo-A-specific sequences that face the extracellular space at the plasma membrane can also be exposed at the cytoplasmic side of intracellular Nogo. This would support the hypothesis that Nogo-A exists in at least two different membrane topologies in oligodendrocytes. The topogenic contribution of the unusually long hydrophobic domains (35 and 36 aa) is thought to be important for this uncommon membrane orientation of the Nogo proteins. It is unclear whether Nogo-A that is present at the cell surface of oligodendrocytes is translocated in the ER in two different orientations or whether it changes its topology at a later stage, e.g., in the Golgi or at the plasma membrane, analogous to certain viral envelope glycoproteins of hepatitis B and C virus (Ostapchuk et al., 1994; Cocquerel et al., 2002).

Nogo-A has no signal sequence. As an alternative, many unconventional membrane proteins can use internal hydrophobic sequences as signal anchor (Goder and Spies, 2001); for Nogo-A this could be one of the C-terminal transmembrane domains. The N-terminal domain would then be translocated only after synthesis has been completed, as has been described for other proteins (Lu et al., 1998; Monne et al., 1999; Nilsson et al., 2000; Goder and Spies, 2001; Cocquerel et al., 2002). It is also possible that Nogo-A reaches the cell surface by a nonconventional pathway, for example by inserting directly into the plasma membrane. For several proteins that have been shown to exhibit more than one topological orientation, such as P-glycoprotein, ductin, cytochrome P450, microsomal epoxide hydrolase, or prion protein, the different topological forms can result in protein targeting to different cellular compartments or correlate with multiple biological functions (for review, see Levy, 1996; Hegde et al., 1998).

Cell-surface Nogo-A comprises ~1% of the total cellular Nogo-A. The large intracellular pool of Nogo-A present in oligodendrocytes may represent an unprocessed precursor form [as for cystic fibrosis transmembrane regulator (CFTR)] (Lu et al., 1998) and/or may have an intracellular, additional function as for, e.g., members of the S100 protein family (for review, see Donato, 1999), CFTR (Bradbury, 1999), and the inositol 1,4,5-trisphosphate receptor (Cunningham et al., 1993; Bush et al., 1994; Mayreiter et al., 1995; Quinton and Dean, 1996; Tanimura et al., 2000).

Additional studies are required to unequivocally demonstrate the herein proposed surface exposure of Nogo-A-specific and Nogo-66 epitopes. Moreover, the identification of the exact topology of the two large C-terminal hydrophobic regions, e.g., by TROSY-based nuclear magnetic resonance (Fernandez et al., 2001), could further clarify our model of two different membrane orientations of Nogo-A.

In summary, our data demonstrate that the Nogo-A-specific part of the molecule has neurite growth inhibitory properties in vitro, and we propose that it is present at the cell surface of certain cells. Different stretches of Nogo-A contribute to this inhibitory activity. The Nogo-A-specific stretches are shown to bind to the surface of responsive cells and brain cortical membranes, suggesting the presence of new, so far unknown, Nogo-A-specific receptor(s) functioning independently of or in a complex with the Nogo-66 receptor NgR. Results from several experiments indicate that Nogo-A can have at least two different membrane topologies, pointing to the possibility of multiple functions both at the cell surface and intracellularly. The data presented here contribute to the understanding of the mechanisms underlying Nogo-mediated neurite growth inhibition and provide important information for the identification of new interacting molecules (receptors, intracellular binding partners) involved in Nogo processing and signaling.

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