An engineered IN-1 F\textsubscript{ab} fragment with improved affinity for the Nogo-A axonal growth inhibitor permits immunochemical detection and shows enhanced neutralizing activity

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The myelin axonal growth inhibitor NI-220/250 (Nogo-A) has attracted considerable attention in elucidating the mechanisms that account for the lack of plasticity in the adult central nervous system. The cognate monoclonal antibody IN-1, which was obtained prior to the molecular characterization of its Nogo-A antigen, has played a crucial role in this respect. However, this murine IgM/k antibody does not only provide an inappropriate format for \textit{in vivo} studies, its low antigen affinity has also hampered the thorough structure-function analysis of its neutralizing effect toward the Nogo-A inhibitor on a molecular basis. We describe here the affinity maturation of a bacterially produced functional IN-1 F\textsubscript{ab} fragment via protein engineering. A soluble fragment of Nogo-A derived from the central exon 3 of its gene, which was prepared by secretion into the periplasm of \textit{Escherichia coli}, served as a target in these experiments. After repeated cycles of site-directed random mutagenesis and screening, the mutant H.1.8 of the IN-1 F\textsubscript{ab} fragment was obtained, carrying five side chain substitutions within CDR-L3. Its dissociation constant for the complex with the recombinant Nogo-A fragment was determined in surface plasmon resonance measurements as approximately 1 \textmu M. The affinity of the unmutated IN-1 F\textsubscript{ab} fragment was 8-fold lower. The engineered F\textsubscript{ab} fragment appeared to be well suited for the specific detection of Nogo-A in immunochemical assays and for the histochemical staining of myelin-rich tissue sections. Most importantly, its concentration-dependent neutralizing effect on the Nogo-A inhibitory activity was significantly enhanced in cell culture. This study confirms Nogo-A to be the antigen of the IN-1 antibody and it demonstrates increased potential of the engineered F\textsubscript{ab} fragment as a reagent for promoting axonal regeneration \textit{in vivo}.

\textbf{Keywords}: affinity maturation/axonal regeneration/bacterial expression/neuronal inhibitor/random mutagenesis

\textbf{Introduction}

The very limited capacity of the adult central nervous system (CNS) for axonal regeneration is a phenomenon of broad and ongoing scientific as well as medical interest (Ramon y Cajal, 1928; Horner and Gage, 2000). In contrast, sprouting and elongation of lesioned axons readily occurs in the peripheral nervous system (PNS). Inhibitory effects and non-permissible properties of CNS tissue, in particular of CNS myelin and oligodendrocytes, probably contribute considerably to the restriction of neuronal regeneration and plasticity. \textit{In vitro}, CNS myelin and oligodendrocyte membranes induce growth cone collapse (Bandtlow \textit{et al.}, 1990).

Based on earlier observations of the inhibitory effect of CNS myelin on neurite outgrowth (Caroni and Schwab, 1988a; Spillmann \textit{et al.}, 1997, 1998) the myelin-associated neurite growth inhibitor NI-220 (Spillmann \textit{et al.}, 1998), later called Nogo-A (Huber and Schwab, 2000), was identified in bovine spinal cord tissue as a predominant protein of oligodendrocytes that prevents axonal growth. The corresponding cDNAs from rat and man were recently described (Chen \textit{et al.}, 2000; GrandPré \textit{et al.}, 2000; Prinjha \textit{et al.}, 2000). The nogo gene encodes three distinct proteins, Nogo-A, Nogo-B and Nogo-C, which apparently arise by alternative splicing and/or promoter usage. Of those, just the full-length Nogo-A transcript is specifically expressed in oligodendrocytes and hence is made mainly responsible for their neuronal growth inhibitory activity (Spillmann \textit{et al.}, 1998; Chen \textit{et al.}, 2000).

The murine monoclonal IgM/k antibody IN-1 was originally raised against the homologous protein NI-250 from rat CNS myelin (Caroni and Schwab, 1988b) and cross-reacts both with the bovine and human protein (Spillmann \textit{et al.}, 1997, 1998). IN-1 was shown to effectively neutralize the inhibitory activity \textit{in vitro} (Bandtlow \textit{et al.}, 1990; Spillmann \textit{et al.}, 1998) and \textit{in vivo}, giving rise to long-distance regeneration and improved plastic changes of injured CNS fibre tracts (Schnell and Schwab, 1990; Z’Graggen \textit{et al.}, 1998). The variable domain cDNAs of this antibody were cloned from the hybridoma cell line, followed by the bacterial production of the corresponding recombinant murine F\textsubscript{ab} fragment, whose functionality was demonstrated \textit{in vitro} (Bandtlow \textit{et al.}, 1996). A partially humanized IN-1 F\textsubscript{ab} fragment was produced by \textit{Escherichia coli} fermentation and shown to successfully promote regeneration of corticospinal axons in adult rats after spinal cord lesion \textit{in vivo} (Broesamle \textit{et al.}, 2000). The recombinant IN-1 F\textsubscript{ab} fragment also induced significant elongation of injured cochlear fibres upon intrathecal treatment (Tatagiba \textit{et al.}, 2002) and a pronounced sprouting response of Purkinje cells after injection into the intact adult cerebellum (Buffo \textit{et al.}, 2000).

Unfortunately, the IN-1 antibody as well as its recombinant F\textsubscript{ab} fragment exhibit rather low antigen affinity, suggesting that the IgM antibody was a product of the early immune response (Bandtlow \textit{et al.}, 1996). Its light chain was found to be encoded by a Vk germine gene devoid of somatic mutations. In the case of the IgM, the poor affinity is partially compensated by an avidity effect due to its decavalent nature, which is attributable to a more general mechanism in immunology, even though at the expense of relaxed specificity in the early humoral response. Consequently, the apparently weaker binding activity of the monovalent F\textsubscript{ab} fragment necessitates higher concentrations in order to provoke comparable \textit{in vitro} activity (Bandtlow \textit{et al.}, 1996).
Nevertheless, in comparison with the intact IgM the use of the recombinant Fab fragment is attractive for several reasons. First, the much smaller Fab fragment was shown to possess high stability in buffer solution and to exert efficient tissue penetration (Bandtlow et al., 1996; Broesamle et al., 2000). Secondly, the Fab fragment can be produced at high yields and quickly purified to homogeneity using a well optimized bacterial expression system (Fiedler and Skerra, 1999, 2001a,b). Finally, its recombinant format permits the application of protein engineering techniques not only to improve its antigen-binding activity but also to humanize the mouse-specific immunoglobulin sequences for medical use.

Here we report the improvement of the functional affinity of the recombinant IN-1 Fab fragment by virtue of introducing a small set of amino acid substitutions into its antigen-binding site. In order to perform cycles of random mutagenesis and selection for enhanced binding activity the presumed extracellular domain of the Nogo-A antigen (Chen et al., 2000; Huber and Schwab, 2000) was produced in a bacterial expression system, thus also permitting molecular characterization of the cognate antigen–antibody interaction for the first time.

Materials and methods

General DNA methodology

DNA manipulations were performed according to standard techniques (Sambrook et al., 1989). Vectors, genes and bacterial strains were from the authors’ collection. DNA polymerases, modifying and restriction enzymes were from generic vendors (New England Biolabs, Frankfurt/Main, Germany; Promega, Mannheim, Germany; Stratagene, Amsterdam, Netherlands). Oligodeoxynucleotides for PCR and DNA sequencing were purchased from Interactiva (Ulm, Germany). PCR reactions were performed using a standard protocol, employing phosphorothioate primers in conjunction with Pfu DNA polymerase (Sambrook, 1992). Site-directed mutagenesis was carried out with single-stranded plasmid DNA following the method of Kunkel et al. (Kunkel et al., 1987) according to a standardized procedure (Geisselsoder et al., 1987). All plasmid constructions and mutagenesis experiments were confirmed by restriction analysis, followed by DNA sequencing of relevant regions on an ABI-Prism™ 310 Genetic Analyzer (Perkin-Elmer/Applied Biosystems, Weiterstadt, Germany) using the BigDye™ terminator kit.

Vector construction for Nogo fragments

A 2.3 kb Nogo-A gene fragment was amplified from the cloned cDNA (Chen et al., 2000) via PCR with the primers 5'-GCT CAG CGG CCG AGA CCC TTT TTG CTC TTC CTP(S)-3' (the Eag I restriction site is underlined) and 5'-GCT TTT AAC TAT GCT GCC CAT TTC TGp(S)T-3'. The single PCR product was digested with Eag I, purified from a 1% agarose gel, and inserted into the multiple cloning region of pASK111 (Vogt and Skerra, 2001), which had been cut with Bsal (resulting in a sticky end compatible with Eag I) as well as Eco47III, yielding pASK111-NiFr1. This vector leads to the production of a mature protein with a molecular mass of 85.0 kDa, including the Strep-tag (Skerra and Schmidt, 2000) at the C-terminus, after processing of the OmpA signal peptide fused in-frame to the N-terminus. pASK111-NiFr2 was constructed from pASK111-NiFr1 by precisely deleting the N-terminal 59 codons from the cloned Nogo-A gene fragment via site-directed mutagenesis using the oligodeoxynucleotide 5'-GAC ATT GAG CTC TTC TAA AAG AGG CCT GCG CTA CGG TAG C-3'. Furthermore, Cys residues were replaced by Ser via site-directed mutagenesis with single-stranded DNA prepared from pASK111-NiFr2 using appropriate oligodeoxynucleotide primers.

The C-terminal Strep-tag encoded on pASK111-NiFr2 was exchanged for a His6 affinity tag by site-directed mutagenesis with the oligodeoxynucleotide 5'-CAC TTC ACA GGTCAA GCT TAT TAA TGG TGA TGG TGA TGG CGT CTT TTA ACT ATG CTG CCC-3'. A KasI restriction site was concomitantly introduced at the 5'-end of the cloned Nogo-A structural gene using the oligodeoxynucleotide 5'-GTT ATC CAT GTT CTT TAA AAG AGG CGG CCT GCC GTT CTA CGG TAG C-3' (the KasI recognition site is underlined), resulting in the vector pASK111-NiFr3. The region encoding the Nogo-A fragment together with the His6-tag was finally subcloned via KasI and NsiI (cutting within the vector, downstream of the Cam' gene) on pASK-IBA4 (Skerra and Schmidt, 2000), which provided the coding region for an N-terminal Strep-tag directly downstream of the OmpA signal sequence. The resulting vector was dubbed pASK111-NiFr4.

Vector construction for Fab fragments

The IN-1 Fab fragment and its mutants were produced utilizing the vectors pASK88, pASK106 or pASK107. All of them encode a chimeric Fab fragment with variable domains derived from the mouse monoclonal antibody IN-1 (Bandtlow et al., 1996) and human constant domains belonging to the subclass IgG1/k (Schiweck and Skerra, 1995). Secrecion into the oxidizing milieu of the bacterial periplasm is ensured by the presence of signal peptides at the N-termini of both chains (Skerra, 1994b) and transcription of the artificial dicistronic operon is under tight control of the chemically inducible tet promoter (Skerra, 1994a). pASK88 (Schiweck and Skerra, 1995) was used for soluble expression and purification via the His6-tag attached to the C-terminus of the heavy chain (Skerra, 1994b; Fiedler and Skerra, 2001a), whereas pASK107 (M.Fiedler and A.Skerra, unpublished results) provided the Strep-tag II for streptavidin affinity purification (Skerra and Schmidt, 2000) instead. pASK110 codes for a Fab fragment similar to pASK88 but with an albumin-binding domain (ABD) appended to the C-terminus of the light chain (König and Skerra, 1998). The variable domain genes were exchanged between the differing vector formats using conserved restriction sites as described (Skerra, 1994b).

Single amino acid exchanges within the IN-1 Fab fragment or its mutants were introduced by site-directed mutagenesis. For this purpose single-stranded DNA of the corresponding vectors pASK88-IN1 or pASK106-L.2.6 was used in conjunction with appropriate oligodeoxynucleotide primers. Random amino acid substitutions were introduced into the variable domain (V\_L) gene of the IN-1 light chain at defined positions via PCR by means of degenerate oligodeoxynucleotide primers (without the phosphorothioate modification) in conjunction with Tag DNA polymerase. Amplification was performed on pASK85-IN1 with the originally cloned genes (Bandtlow et al., 1996) as template. The forward primer 5'-GAC ATT GAG CTC ACC CAG TCT CCA GCA ATC ATG KCT GC-3' (SsrI restriction site underlined) was used in all experiments whereas the oligodeoxynucleotide 5'-GCG CTT CAG CTC GAG CTT GGT CCC AGC TCC GAA CGT MNN AGG MNN MNN TAA CACATT TTG ACA GTA-3' (XhoI restriction site underlined) served as backward primer for randomizing the CDR-L3 positions L93, L94 and L96 at the first stage of
the affinity maturation process. The second mutagenesis cycle was performed with pASK88-L.2.6(153V) as template and the oligodeoxynucleotide 5′-GCC GTT CAT CTC GAG CTT GGT CCC AGC TCC AAC CGG AAC CCG MNN MNN ATT TTG ACA GTA ATC GTG TGC-3′ as second primer for randomizing the positions L91 and L92 together with fixed mutations at L93, L94 and L96. In each case, a single PCR product was obtained, purified from a 1% agarose gel, and cut with SfiI and XhoI. The resulting DNA fragment of approximately 300 bp was ligated with the likewise cut vector backbone of pASK106-IN1 (see above). Colonies of the resulting DNA fragment were grown in 2 l of Luria–Bertani (LB) medium supplemented with the corresponding antibiotic (Am in the case of vectors pASK88, pASK106 and pASK107, or Cam for pASK110) at 22°C and 200 r.p.m. Gene expression was induced by application of a 1% (w/v) L-arabinose solution at an optical density (OD) of 0.5 at 550 nm by addition of 0.46 mg/ml anhydrotetracycline (aTc; Acros Organics, Geel, Belgium) in the case of the Fab, fragments or of 400 µg/1 µl aTc in the case of Nogo-A fragments. After 3 h induction the bacteria were harvested by centrifugation and the periplasmic protein fraction was prepared as described (Skerra and Schmidt, 2000). Bacterial periplasm was recovered by means of the His 6-tag via immobilized metal affinity chromatography (IMAC) (Skerra, 1994b) using 50 mM NaPi, pH 7.5, 1 M NaCl as the chromatography buffer. The elution was effected by application of a linear gradient of 0 to 75 mM imidazole in the same buffer. The specific enzyme activities of the proteins were determined using calculated absorption coefficients at 280 nm (Gill and von Hippel, 1989) of 0.41 ml mg⁻¹ cm⁻¹ for the Nogo-A fragments and of 1.8 ml mg⁻¹ cm⁻¹ for the IN-1 Fab fragments, respectively.

Bacterial protein production

Cultures of E. coli JM83 transformed with the appropriate expression vector were grown in 2 l of Luria–Bertani (LB) medium supplemented with the corresponding antibiotic (Am for derivatives of vectors pASK88, pASK106 and pASK107, or Cam for pASK110) at 22°C and 200 r.p.m. Gene expression was induced by application of a 1% (w/v) L-arabinose solution at an optical density (OD) of 0.5 at 550 nm by addition of 0.46 mg/ml anhydrotetracycline (aTc; Acros Organics, Geel, Belgium) in the case of the Fab, fragments or of 400 µg/1 µl aTc in the case of Nogo-A fragments. After 3 h induction the bacteria were harvested by centrifugation and the periplasmic protein fraction was prepared as described (Skerra and Schmidt, 2000). For the production of Nogo-A fragments 200 µg/ml lysozyme was added to the cell fractionation buffer (50 mM NaP, pH 7.5, 500 mM sucrose, 1 mM EDTA).

Nogo-A fragments NI-Fr1 and NI-Fr2 were purified from the periplasmic protein extract via the Strep-tag fused to their C-termini employing streptavidin affinity chromatography (Skerra and Schmidt, 2000), whereby elution was achieved under mild conditions in the presence of D-desiobiotin. After dialysis against a chromatography buffer (50 mM NaP, pH 7.5, 150 mM NaCl, 1 mM EDTA) and the concentration of the eluate (Vivaspin 15, MWCO 30 kDa; Greiner, Frickenhausen, Germany), further purification was achieved by gel filtration on a Superdex™ 200 prep grade column (Pharmacia, Uppsala, Sweden) using Dynamax™ SD-300 HPLC equipment (Rainin, Woburn, MA). NI-Fr4 was first purified by means of the His 6-tag via immobilized metal affinity chromatography (IMAC) (Skerra, 1994b) using 50 mM NaP, pH 7.5, 1 M NaCl as the chromatography buffer and a linear elution gradient from 0 to 75 mM imidazole-HCl in the same buffer. The specifically eluted protein fraction was then subjected to streptavidin affinity chromatography as above.

The recombinant IN-1 Fab fragments were purified either by IMAC via the His 6-tag fused to the C-terminus of their heavy chain (Fiedler and Skerra, 2001a) or, when using pASK107 (see above), via streptavidin affinity chromatography (Schlapschy and Skerra, 2001). IMAC was also performed under FPLC conditions using a POROS MC/M column (0.46×10 cm; PerSeptive Biosystems, Wiesbaden, Germany) charged with Zn²⁺ ions and Dynamax™ SD-300 HPLC equipment (Rainin) operating at a flow rate of 2 ml/min. A 12.5 ml aliquot of periplasmic extract from a 2 l E. coli culture dialysed against 50 mM NaP, pH 7.5, 500 mM betaine was applied to the column and, after washing with dialysis buffer, elution was effected by application of a linear gradient of 200 mM imidazole-HCl, pH 7.5, 50 mM NaP, 500 mM betaine against dialysis buffer. This method enabled a 5-fold quicker purification compared with the conventional procedure (Fiedler and Skerra, 2001a), yielding recombinant Fab fragments with an apparent purity of >95% as estimated from SDS–PAGE.

The yields of purified recombinant proteins from 2 l shaker-flask experiments were highly reproducible but varied between 0.04 and 0.8 mg 1⁻¹ OD⁻¹ for the different Fab fragments and between 0.1 and 0.3 mg 1⁻¹ OD⁻¹ for the Nogo-A fragments. After purification the proteins were stored in PBS (4 mM KH₂PO₄, 16 mM Na₂HPO₄, 115 mM NaCl) containing 0.1 mM EDTA at 4°C for up to several weeks. Protein purity was checked by SDS–PAGE using 0.1% (w/v) SDS, 10 or 15% (w/v) polyacrylamide gels (Fling and Gregerson, 1986) stained with Coomassie brilliant blue. The concentration of the purified recombinant proteins was determined using calculated absorption coefficients at 280 nm (Gill and von Hippel, 1989) of 0.41 ml mg⁻¹ cm⁻¹ for the Nogo-A fragments and of 1.8 ml mg⁻¹ cm⁻¹ for the IN-1 Fab fragments, respectively.

Filter-sandwich colony-screening assay

This assay was carried out based on published procedures (Skerra et al., 1991; Schlehuber et al., 2000). Transformed E. coli JM83 cells harbouring the pASK106 vector encoding Fab fragments fused with the ABD (see above) were plated on a hydrophilic membrane (GVWP, 0.22 µm; Millipore, Bedford, MA), placed on a Petri dish with LB/Amp agar, such that approximately 500 colonies were obtained, and incubated at 37°C for 8–9 h. In the meantime, a hydrophobic membrane (Immobilon™-P, 0.45 µm; Millipore) was coated with 10 mg/ml human serum albumin (HSA; Sigma, Deisenhofen, Germany) in PBS for 4 h and blocked with 3% (w/v) BSA (Roth, Karlsruhe, Germany), 0.5% (v/v) Tween-20 in PBS. The membrane was washed twice with PBS, soaked in LB/Amp containing 200 µg/ml aTc, and placed on an LB/Amp agar plate supplemented with 200 µg/ml aTc. The first membrane, carrying tiny colonies of the transformed cells, was then placed onto the second (hydrophobic) membrane. The filter sandwich was incubated for 16 h at 22°C. During this period the mutated IN-1 Fab fragments were secreted—and partially released from the colonies by leakage from the bacterial periplasm—and finally immobilized on the lower membrane via complex formation between HSA and ABD.

The first membrane with the still viable colonies was transferred to a fresh LB/Amp agar plate and stored at 4°C. The second membrane was washed three times in PBS containing 0.1% (v/v) Tween-20 (PBS/T) and the immobilized Fab fragments, each in a spot corresponding to the position of the original colony, were probed for antigen binding. To this end recombinant Nogo-A fragment was labelled at a molar ratio of 5:1 with digoxigenin-3-O-methylcarbonyl-ε-aminocaproic acid N-hydroxy-succinimide ester (Roche Diagnostics, Mannheim, Germany) and applied to the membrane for 1 h at a concentration of 30 or 50 µg/ml in PBS/T. After washing three times with PBS/T the membrane was incubated for 1 h with 0.75 U/ml anti-digoxigenin Fab fragment conjugated with alkaline phosphatase (Roche Diagnostics) in 10 ml of PBS/T. The membrane was finally washed twice with PBS/T and twice with PBS and the signals were developed using standard chromogenic substrates as described (Schlehuber et al., 2000). Colonies corresponding to signals with an intensity above average were identified, recovered from the first membrane, and propagated for further analysis of their recombinant gene products.
Enzyme-linked immunosorbent assay (ELISA)

ELISA was carried out in a 96-well microtitre plate (Becton Dickinson, Heidelberg, Germany) at ambient temperature with incubation steps of 1 h unless otherwise stated. Three washing steps with PBS/T were used after each incubation, and residual liquid was removed thoroughly. The wells were coated for 4 h with 50 µl of a solution of NI-Fr2 at concentrations between 180 and 200 µg/ml in PBS buffer and then blocked with 200 µl of 3% (w/v) BSA, 0.5% (v/v) Tween-20 in PBS. After washing, 50 µl of the purified recombinant Fab fragment was applied as a dilution series in PBS/T. The wells were then incubated with 50 µl of anti-human C_\kappa antibody conjugated with alkaline phosphatase (Sigma), diluted 1:1000 in PBS/T. Signals were finally developed in the presence of p-nitrophenyl phosphate (Voss and Skerra, 1997). Enzymatic activity was measured at 25°C as the change in absorbance at 405 nm/min with a SpectraMAX 250 instrument (Molecular Devices, Sunnyvale, CA). The data were corrected for background values determined in wells that were merely coated with BSA and fitted by non-linear least squares regression as described (Voss and Skerra, 1997).

Real-time surface plasmon resonance (SPR) measurements

Molecular interaction analysis was performed using a Biacore-X® system equipped with the NTA-sensor chip® (Biacore AB, Uppsala, Sweden). PBS containing 0.005% (v/v) surfactant P20 was used as continuous flow buffer as well as for dilution of proteins. Analysis was performed at 25°C using a flow rate of 35 µl/min. For each measurement the derivatized chip surface was charged with 70 µl of 0.5 mM NiSO_4, followed by immobilization of NI-Fr4 via its His_\_tag in one of the two flow channels by applying 70 µl of a 50 µg/ml solution of the purified recombinant protein. Then, the Fab fragment (produced by means of the vector pASK107 and purified via the Strep-tag II) was injected at a de

Cryosections (12 µm) of rat brain (Rattus norvegicus) were fixed for 10 min using ice-cold ethanol. The following incubation steps were then each performed for 1 h at room temperature in a humid chamber using PBS. Unless otherwise stated, slides were washed for 5 min with PBS. After blocking with 4% (w/v) BSA, the Fab fragment (produced using the pASK88 vector type and purified via the His_\_tag) was applied at a concentration of 100 µg/ml. After three washing steps, bound Fab fragment was detected with an anti-human C_\kappa antibody alkaline phosphatase conjugate (Sigma), diluted 1:100. The sections were then washed three times with TBS (25 mM Tris–HCl, pH 7.4, 145 mM NaCl, 3 mM KCl) and staining was performed using the Fast red kit (Roche Diagnostics). The microscopic slides were photographed on an Axiohot microscope (Carl Zeiss, Jena, Germany) using 20-fold magnification.

Neurite outgrowth assay

Neurite growth-modulating properties of the different Fab fragments were tested on 4-well plastic dishes (Greiner, Nürtlingen, Germany) coated with recombinant Nogo-A. Briefly, test wells were coated for 20 min with 100 µg/ml poly-L-lysine, washed with Hank’s balanced salt solution (HBSS; Life Technologies, Basel, Switzerland) and coated for 2 h with 15 or 30 µg/ml of recombinant rat Nogo-A (entire extracellular domain refolded from bacterial inclusion bodies; T.Oertle et al., manuscript in preparation). Recombinant Nogo-A was omitted in the wells serving for control. After aspiration, the wells were washed with Dulbecco’s modified Eagle’s medium (Life Technologies) containing 10% (v/v) fetal calf serum (Life Technologies) and blocked in the same medium for 20 min at 37°C.

Cerebellar cell cultures were prepared from rat on postnatal day 7/8. Cells were dissociated by combined trituration and trypsinization and purified on Percoll gradients as described (Hatten, 1985). The cerebellar granule cells were plated in chemically defined neurobasal medium supplemented with B27 and 0.2 mM glutamine, 100 U/ml penicillin, and 0.1 mg/ml streptomycin (Life Technologies). To assess the neutralization of inhibitory activity, substrate-coated wells were first incubated with 100 µg/ml of the different recombinant Fab fragments dialysed against NaCl/P_6 (137 mM NaCl, 2.7 mM KCl, 1.5 mM KH_2PO_4, 8 mM Na_2HPO_4, pH 7.4) for 20 min at 37°C. The wells were then washed briefly with HBSS and cells were applied in the presence of the Fab fragments.

Assays were stopped after 24 h in culture by adding 4% (v/v) formalin buffered with NaCl/P_6. For assaying the inhibitory substrate properties, the proportion of total cells bearing neurites longer than the diameter of the cell body (indicating that neurite outgrowth was successfully initiated) was determined. Under control conditions, i.e. in the absence of recombinant Nogo-A, 70% of the cerebellar granule neurons formed processes. Quantification of neurite lengths was performed on cultures monitored with a Zeiss Axiohot microscope. Phase contrast pictures were acquired with a 12-bit digital CCD camera (Visicam Visitor, Germany) and analysed using Metamorph software (Universal Imaging Corporation, West Chester, PA). For each well, the longest neurites of at least 100 isolated neurons were measured and averaged. Three wells were investigated for each experimental condition.

Results

Bacterial synthesis of a soluble Nogo-A domain

Although the precise transmembrane topology of the Nogo-A protein has yet to be experimentally determined, its primary structure—as deduced from the cDNA sequence (Chen et al., 2000)—reveals several characteristic features (Figure 1). The rat full-length protein comprises 1163 amino acids. Its N-terminal region does not reveal a secretory signal sequence as would be expected for a normal class I membrane protein (von Heijne and Gavel, 1988). Instead it is highly negatively charged. In particular, the amino acid stretch from positions 31 to 50 is almost entirely (with the exception of a single Pro) composed of Glu and Asp residues. The acidic sequence is followed by a region rich in Pro, Ala and— to a lesser extent— also Ser and Gly residues, which reaches from position 61 to 171, i.e. directly up to the end of exon 1, which is followed by a very small exon 2 (T.Oertle et al., manuscript in...
IN-1 Fab fragments with improved affinity for Nogo-A

**Fig. 1.** Recombinant Nogo-A fragments prepared in this study. (A) Structural characteristics of the native neurite growth inhibitor Nogo-A and of the recombinant fragments derived from it. (B) SDS–PAGE analysis of bacterially produced NI-Fr4. Periplasmic protein extract from *E. coli* JM83 harbouring pASK111-NIFr4 (lane 1); flow-through of IMAC column (lane 2); eluted protein from IMAC column as applied to the streptavidin column (lane 3); flow-through of streptavidin column (lane 4); purified protein after streptavidin affinity chromatography (lane 5). Molecular sizes are indicated at the left.

preparation). Thus, the domain encoded by exon 1 is likely not to adopt a regular secondary structure. In contrast, the large Nogo-A-specific exon 3 reveals an average amino acid composition similar to that of globular proteins. Furthermore, it exhibits altogether eight Cys codons, whereas otherwise this amino acid is absent in the full-length protein. This feature might point towards the presence of disulphide bonds in the exon 3 region as they are frequently found in extracellular domains of membrane proteins. The C-terminal domain encoded by exons 4–9 (for the precise exon structure refer to T.Oertle, *et al.*, manuscript in preparation) is short again, comprising altogether 188 residues. It includes two stretches of 36 (residues 990–1025) and 35 (residues 1092–1126) amino acids, respectively, with a high content of hydrophobic side chains, thus probably constituting transmembrane regions.

From the structural point of view the protein fragment encoded by exon 3 is most likely folded as a globular domain and should carry the biological active site. Consequently, we chose the polypeptide comprising residues 174–940 (containing 767 residues, i.e. 66% of full-length Nogo-A) for production as a recombinant protein. The gene fragment was amplified from the cDNA and subcloned on a standard vector for protein production in *E. coli*. Using appropriate PCR primers, the Nogo-A fragment was precisely fused at its N-terminus (i.e. in front of residue 174) to the OmpA signal peptide, thus effecting secretion into the bacterial periplasm, where efficient disulphide bond formation is favoured by an oxidizing redox environment. At the C-terminus (i.e. following residue 940) the fragment was fused with the Strep-tag affinity peptide, conferring binding activity towards streptavidin for simplified purification (Skerra and Schmidt, 2000). Transcription of the resulting hybrid gene was under tight control of the tetracycline promoter/operator (Skerra, 1994a). The resulting protein was denominated NI-Fr1 (Figure 1A).

Upon induction of gene expression NI-Fr1 was readily liberated from the periplasmic protein fraction of *E. coli* and purified by streptavidin affinity chromatography in one step. SDS–PAGE analysis revealed that approximately 50% of the recombinant protein comprised a product with the proper length whereas 50% corresponded to a series of smaller polypeptides, probably representing proteolytic degradation products (data not shown). In particular, there appeared one
prominent band just underneath that for the major recombinant protein. Both bands were subjected to N-terminal sequencing. The upper band yielded the sequence Glu–Thr–Leu–Phe–Ala, which resulted from the precise cleavage of the OmpA signal peptide. The lower band started with the amino acids Ser–Phe–Lys–Glu–His, i.e. at a position 59 codons downstream within the cloned sequence (beginning at residue 233 in the full-length primary structure). Its appearance was most likely due to the action of a bacterial protease and might indicate that the N-terminal part of the chosen Nogo-A fragment still belongs to a polypeptide segment devoid of well defined structure.

In order to achieve better homogeneity of the gene product, the first 59 residues of the mature polypeptide chain were deleted from the cloned coding region, leading to NI-Fr2 (Figure 1A). This protein was readily produced in the periplasm of E.coli, with similar yields as the former version but clearly a reduced degradation pattern. The possible presence of structural disulphide bonds in the recombinant protein was investigated by individually substituting all eight Cys residues (corresponding to positions 323, 403, 443, 536, 574, 676, 885 and 890 in the full-length Nogo-A sequence) with Ser via site-directed mutagenesis. The eight mutant Nogo-A fragments were produced in E.coli as before. However, it was not possible to recover the mutants Cys323→Ser and Cys885→Ser from the periplasmic protein fraction, while the mutants Cys443→Ser and Cys890→Ser gave rise to significantly diminished yields after Strep-tag purification when compared with the wild-type protein. In contrast, the other four mutants were produced at similar amounts as the original Nogo-A fragment. These observations indicate that at least some of the Cys residues are important for folding and may be involved in cystine crosslinks.

The wild-type NI-Fr2 protein still gave rise to certain truncated products, which was considered undesirable for precise binding measurements (see below). Notably, for the bovine bNI-220 susceptibility towards proteolysis had been investigated by individually substituting all eight Cys residues (corresponding to positions 323, 403, 433, 536, 571, 674, 885 and 890 in the full-length Nogo-A sequence) with Ser via site-directed mutagenesis. The eight mutant Nogo-A fragments were produced in E.coli as before. However, it was not possible to recover the mutants Cys323→Ser and Cys885→Ser from the periplasmic protein fraction, while the mutants Cys443→Ser and Cys890→Ser gave rise to significantly diminished yields after Strep-tag purification when compared with the wild-type protein. In contrast, the other four mutants were produced at similar amounts as the original Nogo-A fragment. These observations indicate that at least some of the Cys residues are important for folding and may be involved in cystine crosslinks.

The wild-type NI-Fr2 protein still gave rise to certain truncated products, which was considered undesirable for precise binding measurements (see below). Notably, for the bovine bNI-220 susceptibility towards proteolysis had been observed before (Spillmann et al., 1998). Consequently, a doubly tagged version of the recombinant protein was prepared using an otherwise identical expression system. First, the Strep-tag at the C-terminus was exchanged for a His_6-tag (yielding NI-Fr2, Figure 1A). This protein was readily produced in the periplasm of E.coli, with similar yields as the former version but clearly a reduced degradation pattern. The possible presence of structural disulphide bonds in the recombinant protein was investigated by individually substituting all eight Cys residues (corresponding to positions 323, 403, 443, 536, 574, 676, 885 and 890 in the full-length Nogo-A sequence) with Ser via site-directed mutagenesis. The eight mutant Nogo-A fragments were produced in E.coli as before. However, it was not possible to recover the mutants Cys323→Ser and Cys885→Ser from the periplasmic protein fraction, while the mutants Cys443→Ser and Cys890→Ser gave rise to significantly diminished yields after Strep-tag purification when compared with the wild-type protein. In contrast, the other four mutants were produced at similar amounts as the original Nogo-A fragment. These observations indicate that at least some of the Cys residues are important for folding and may be involved in cystine crosslinks.

In vitro affinity maturation of the IN-1 Fab fragment

Initial experiments on the detection of natural Nogo-A as well as recombinant Nogo-A fragments on western blots or on tissue sections by means of the bacterially produced IN-1 Fab fragment revealed relatively weak signals (Bandtlow et al., 1996), indicating that the antigen affinity was poor. From the cloning of the variable domain genes of the IN-1 monoclonal antibody and analysis of their sequences it was evident that the VH domain had undergone somatic hypermutation while the VL domain appeared to be a newly defined mouse germline gene (Bandtlow et al., 1996). A computer modelling study was carried out based on a human anti-thyroid peroxidase autoantibody (Chacko et al., 1996; PDB entry 1VGE) and a murine anti-phenylsorionate antibody (Rose et al., 1990; PDB entry 6FAB), both of which have sets of complementarity-determining regions (CDRs) with the same lengths and canonical structure determinants (Chothia et al., 1989) as IN-1 and share a high amino acid sequence similarity with it. This analysis suggested that the CDR-L3 of IN-1 and, to a lesser extent, its CDR-L1, were the most promising target regions for protein engineering towards improved antigen recognition. In particular, residue L96 and also residue L32 (in CDR-L1) were expected to be exposed close to the centre of the combining site and thus likely to be involved in contacts with the antigen.

Within CDR-L1 both IN-1 and 1VGE have an Ala residue at position L32 whereas 6FAB carries a Phe. On the other hand, IN-1 as well as 6FAB carry an Arg at position L96 (in CDR-L3) while 1VGE exhibits a Leu. Therefore, the structural consequences of the amino acid exchanges L^32→F and R^L96→L within the VH domain of IN-1 were modelled, resulting in their identification as potential paratope residues. The corresponding single amino acid exchanges in the recombinant Fab fragment were introduced by site-directed mutagenesis (Fiedler and Skerra, 1999), followed by production in E.coli and purification via IMAC (Fiedler and Skerra, 2001a). A test for neutralizing biological activity in the 3T3 fibroblast assay for inhibition of cell spreading on a CNS myelin substrate (Bandtlow et al., 1996) revealed that the mutant R^L96→L had a slightly improved activity. In contrast, the mutant L^32→F had mostly lost its neutralizing activity when compared with the wild-type IN-1 Fab fragment (data not shown). Therefore, the IN-1(F^L32F) Fab fragment was used as a negative control during in vivo regeneration experiments of lesioned corticospinal tract fibres in rats (Broesamle et al., 2000).

In order to perform functionally more complex changes within the paratype of the IN-1 antibody a cluster of three amino acids in CDR-L3 corresponding to positions L93, L94 and L96, was now subjected to targeted random mutagenesis. All 20 side chains were allowed for substitution in each position, followed by screening for improved binding of the recombinant Nogo-A fragment via a filter-sandwich colony-screening assay (Skerra et al., 1991; Schlehuber et al., 2000). For this purpose, a genetic random library was prepared by PCR amplification of the IN-1 VL gene using a degenerate primer that carried the corresponding mixed base positions (see Materials and methods). The mutated gene fragment was recloned on the expression vector pASK106-IN1 (encoding a Fab fragment fused with an ABD to the C-terminus of its light chain; König and Skerra, 1998). E. coli JM83 was transformed with the ligation mixture and the cell suspension was plated on four filter membranes, placed on top of agar plates, thus screening approximately 2000 colonies in parallel.

The antigen-binding activity of the bacterially secreted mutants of the IN-1 Fab fragment, which became functionally immobilized on a second membrane soaked with HSA (see Materials and methods), was probed by incubation with digoxigenin-labelled NI-Fr2, followed by detection with an anti-digoxigenin Fab fragment conjugated with alkaline phosphatase. Colonies that gave rise to staining signals above average were recovered, propagated, and their plasmids were isolated for DNA sequence analysis. Out of 31 clones that were investigated, 12 plasmids were identified carrying functional VL genes (for the mutations see Table I). Otherwise, frameshift
The substitutions were introduced by site-directed mutagenesis and all corresponding recombinant F\textsubscript{ab} fragments were produced and purified as before, resulting in yields that were similar again to the wild-type IN-1 F\textsubscript{ab} fragment. However, when binding activity towards the recombinant NI-Fr2 antigen was tested in an ELISA, all these mutants gave rise to significantly lower signals than the original I.2.6 F\textsubscript{ab} fragment. Merely the replacement Cys\textsubscript{L96}→Val resulted in a detectable binding behaviour (Figure 2) and was therefore used as the basis for the second affinity maturation cycle.

CDR-L3 forms a connecting loop between two neighbouring β-strands such that the positions L91 and L92 are in close spatial proximity with L96. Hence, in order to structurally compensate a possible misfit at position L96—due to the exchange of Cys by Val—the positions L91 and L92 within CDR-L3 of the I.2.6(F\textsubscript{L96V}) F\textsubscript{ab} fragment were subjected to targeted random mutagenesis and the filter-sandwich colony-screening assay was performed again. This time the stringency of selection was raised by lowering the concentration of the recombinant antigen (a mutant of NI-Fr2 devoid of Cys\textsubscript{574} and Cys\textsubscript{676}) from 50 to 30 µg/ml. From screening approximately 1000 colonies spread on two filter membranes, 16 clones were identified according to their pronounced colour signals. In contrast with the previous experiment, all of them carried plasmids encoding functional mutants of the I.2.6(F\textsubscript{L96V}) F\textsubscript{ab} fragment. The VL gene cassettes of these four clones (Table I) were subcloned on pASK88-IN1 and the corresponding F\textsubscript{ab} fragments were produced and purified as before. One of them, the II.1.8 F\textsubscript{ab} fragment (Figure 2A), exhibited clearly improved binding activity over the I.2.6(F\textsubscript{L96V}) mutant in an ELISA (Figure 2B), even though its affinity was still lower than that of the original I.2.6 mutant carrying the free Cys residue. Nevertheless, the yield of the II.1.8 mutant was 12-fold higher upon expression in E.coli and thus close to that of the recombinant wild-type IN-1 F\textsubscript{ab} fragment (0.5 versus 0.8 mg l\textsuperscript{-1} versus 0.8 mg l\textsuperscript{-1}, respectively).

**Functional analysis of engineered F\textsubscript{ab} fragments**

For a detailed analysis of the antigen-binding activity and application in immunohistochemistry as well as cell culture

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**Table I. Mutants obtained from affinity maturation of the IN-1 F\textsubscript{ab} fragment**

<table>
<thead>
<tr>
<th>Position</th>
<th>Signal in CSA\textsuperscript{a}</th>
<th>Expression yield\textsuperscript{b}</th>
<th>ELISA signal</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L91</td>
<td>L92</td>
<td>L93</td>
</tr>
<tr>
<td>IN-1 wt</td>
<td>Val</td>
<td>Leu</td>
<td>Ser</td>
</tr>
<tr>
<td>I.1.4</td>
<td>–</td>
<td>–</td>
<td>Pro</td>
</tr>
<tr>
<td>I.1.6</td>
<td>–</td>
<td>–</td>
<td>Asn</td>
</tr>
<tr>
<td>I.1.11</td>
<td>–</td>
<td>+</td>
<td>Tyr</td>
</tr>
<tr>
<td>I.1.16</td>
<td>–</td>
<td>+</td>
<td>Met</td>
</tr>
<tr>
<td>I.2.2</td>
<td>–</td>
<td>–</td>
<td>Arg</td>
</tr>
<tr>
<td>I.2.4</td>
<td>–</td>
<td>–</td>
<td>Gly</td>
</tr>
<tr>
<td>I.2.5</td>
<td>–</td>
<td>–</td>
<td>Arg</td>
</tr>
<tr>
<td>I.2.6</td>
<td>–</td>
<td>–</td>
<td>Tyr</td>
</tr>
<tr>
<td>I.2.8</td>
<td>–</td>
<td>–</td>
<td>Arg</td>
</tr>
<tr>
<td>I.2.9</td>
<td>–</td>
<td>–</td>
<td>Phe</td>
</tr>
<tr>
<td>I.4.4</td>
<td>–</td>
<td>–</td>
<td>Asp</td>
</tr>
<tr>
<td>I.2.6(F\textsubscript{L96V})</td>
<td>–</td>
<td>–</td>
<td>Arg</td>
</tr>
<tr>
<td>II.1.1</td>
<td>Arg</td>
<td>Lys</td>
<td>Arg</td>
</tr>
<tr>
<td>II.1.3</td>
<td>Met</td>
<td>Lys</td>
<td>Arg</td>
</tr>
<tr>
<td>II.1.7</td>
<td>Leu</td>
<td>Lys</td>
<td>Arg</td>
</tr>
<tr>
<td>II.1.8</td>
<td>Ile</td>
<td>Asn</td>
<td>Arg</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Filter-sandwich colony-screening assay.

\textsuperscript{b}In E.coli JM83 using the vector pASK88.

\textsuperscript{c}No exchange.
assays, the different engineered versions of the IN-1 F\textsubscript{ab} fragment were produced in \textit{E. coli} in shaker flask cultures and purified by IMAC to homogeneity (Figure 3A). The thermodynamic affinity for the recombinant Nogo-A fragment NI-Fr4 was determined both for the II.1.8 mutant and for the wild-type IN-1 F\textsubscript{ab} fragment using the method of real-time SPR on a Biacore-X\textsuperscript{®} system. Preliminary experiments revealed considerable difficulties in immobilizing the recombinant Nogo-A fragments NI-Fr2 as well as NI-Fr4 to sensor chips carrying the conventional carboxy-methylated dextran matrix (CM5 sensor chip\textsuperscript{®}) via the standard amine coupling procedure (O’Shannessy \textit{et al.}, 1992). Most likely, the extremely low pI of the Nogo-A fragment (calculated to be 4.2 for NI-Fr2 using
the GCG program package; Devereux et al., 1984) led to electrostatic repulsion within the acceptable range of pH conditions.

Therefore, a Ni/NTA-derivatized sensor chip (NTA-sensor chip®; Nieba et al., 1997) was chosen, to which NI-Fr4 was immobilized via its C-terminal His6-tag (Figure 1A; see Materials and methods). In order to prevent binding of the recombinant Fαb fragments to the same surface, their variable domain genes were subcloned on the vector pASK107, thus permitting purification—with comparable efficiency—via the Strep-tag II instead of the His6-tag (see Materials and methods). The Fαb fragments were applied to the sensor chip, each time charged with fresh antigen, at different concentrations and the amount of bound Ig fragment was measured under equilibrium conditions. In this way, binding isotherms were obtained for the wild-type and engineered Fαb fragments (Figure 3B), from which dissociation constants were deduced. The $K_D$ value for the recombinant wild-type IN-1 Fαb fragment was 7.8 ± 1.9 µM. In contrast, the dissociation constant for its II.1.8 mutant was 1.04 ± 0.18 µM, i.e. 8-fold better. Control experiments with an unrelated protein, recombinant cystatin (Auerswald et al., 1989) carrying a His6-tag (A. Skerra, unpublished results), that was used instead of the Nogo-A fragment for coating of the sensor chip, confirmed the absence of unspecific binding (data not shown).

The engineered II.1.8 Fαb fragment was also employed for the detection of natural Nogo-A by immunohistochemistry. Figure 4 shows cross-sections of adult rat brain which were stained with different recombinant Fαb fragments, followed by a secondary antibody conjugated with a reporter enzyme. The II.1.8 mutant specifically stained the myelinated regions, especially the Corpus callosum and transected fibre bundles of the Capsula interna in the Corpus striatum. The staining pattern is similar in morphology and intensity to the one obtained with a recombinant Fαb fragment derived from the monoclonal antibody 8-18C5, which is directed against the major oligodendrocyte glycoprotein MOG (Linington et al., 1984). The staining with the recombinant wild-type IN-1 Fαb fragment was very weak under the present conditions of fixation. An unrelated recombinant anti-CD30 Fαb fragment derived from the HRS-3 antibody (Engert et al., 1990) gave only background staining. These results demonstrate that the affinity of the II.1.8 mutant of the IN-1 Fαb fragment has been raised to a sufficient extent in order to detect the Nogo-A antigen in standard immunohistochemical experiments. Analogous results were obtained using immunofluorescence microscopy (data not shown).

Finally, the engineered Fαb fragments were tested for their neutralizing effect on Nogo-A substrate properties using a cell culture assay. As shown in Figure 5, neurite outgrowth of cerebellar granule cells was severely reduced when recombinant Nogo-A was used as a substrate. In contrast, in its absence, poly-L-lysine promoted extensive attachment of granule cells, as well as robust neurite growth with an average neurite length of approximately 70 µm in 70% of adherent cells. In this in vitro bioassay functional neutralization of the inhibitory Nogo-A substrate was observed at different degrees for the various engineered Fαb fragments (Figure 5). While the recombinant wild-type IN-1 Fαb fragment revealed partial neutralization of Nogo-A activity, as previously demonstrated (Bandtlow et al., 1996), introduction of the mutation AlaL32→Phe into the VL domain completely abolished this effect. In contrast, the mutants 1.2.6(L96V) and, in particular,
II.1.8, exhibited significantly stronger neutralizing effects, as
revealed by their better fibre growth-promoting activities, even
when the concentration of the inhibitory material was raised.
None of the applied F_{ab} fragments exerted an effect on neurite
growth of cerebellar granule cells under control conditions,
i.e. in the absence of Nogo-A. Notably, the stepwise improve-
ment of the biological activity of the mutants I.2.6(L96V) and
II.1.8 in comparison with the wild-type IN-1 F_{ab} fragment
correlated nicely with their relative increase in antigen affinity
observed in the ELISA experiment (Figure 2B).

Discussion

In spite of its considerable potential as a reagent for studying
the biological role and distribution of the Nogo-A antigen and
for neutralizing its inhibitory effect on axonal growth in tissue
culture as well as animal experiments, the use of the monoclonal
antibody IN-1 has so far been hampered by its apparently low
antigen affinity. In the present study we were able to quantify
the intrinsic affinity of its recombinant F_{ab} fragment to be in
the 10^{5} M^{-1} range. The remarkably high dissociation constant
that was measured between the IN-1 F_{ab} fragment and the
bacterially produced Nogo-A domain explains several observa-
tions from previous studies.

First, the antibody fragment had to be applied at a relatively
high concentration, typically 5 mg/ml (corresponding to
approximately 100 µM), in order to provoke a biological effect
(Bandtlow et al., 1996; Broesamle et al., 2000). The original
monoclonal antibody was similarly effective at a 5–10-fold
lower concentration, which is in agreement with what has to
be expected from the strong avidity effect of an IgM as soon
as it interacts with an immobilized antigen at a high surface
density. Unfortunately, the precise concentration of the IgM
was rarely used as an experimental parameter because of the
difficulties associated with its purification from the hybridoma
supernatant.

Secondly, the use of IN-1 in immunochemical detection
experiments, like western blotting, immunohistochemistry,
fluorescence microscopy and immunoprecipitation, was
limited, and the original recombinant F_{ab} fragment was
even dissatisfactory in this respect. In such experiments an
initial incubation period allowing for antibody–antigen
association is usually followed by several washing steps,
where the Ig protein (or the solubilized antigen) may be
quickly lost by a bleeding effect if the kinetics of dissociation
are fast. This usually happens if the thermodynamic dissociation
constant is high and it was also observed here in the course
of our Biacore measurements (data not shown). Unfortunately,
as a general implication of reaction kinetics this phenomenon
cannot be overcome by simply raising the concentration of the
applied complexing partner.

The fundamental nature of this problem explains why it
took an increased effort and repeated cycles of mutagenesis
in order to finally engineer a derivative of the IN-1 F_{ab}
fragment with 8-fold enhanced affinity. The II.1.8 mutant
possesses a K_D value equal to 1 µM, which is obviously
sufficient to yield clear antigen-binding signals in an ELISA
and to effect specific staining in immunohistochemistry on
brain sections. The preceding mutant I.2.6 might actually
perform even better, yet its exposed Cys residue at position
L96 within CDR-L3, which apparently plays a crucial role,
prevents its practical application due to poor expression in
E.coli. In this respect, one might consider switching to another
host organism. However, unpaired exposed Cys residues have
a general tendency of being post-translationally modified in
eukaryotic cells (Lyons et al., 1990) and they provide a
problem during storage because of susceptibility towards
oxidation. Nevertheless, the mutant II.1.8 IN-1 F_{ab} fragment
with its well defined antigen-binding activity presents a promis-
ing starting point for future affinity maturation attempts, where
the mutagenesis could be focused at hypervariable regions
apart from CDR-L3, possibly even within the VH domain.

The present investigation also sheds light on aspects of the
structure and biological function of the Nogo-A antigen. First,
our data provide evidence that the epitope for the IN-1 antibody
(or its F_{ab} fragment, respectively) resides within the protein
domain encoded by exon 3. This exon is exclusively present
in the splice form called Nogo-A, which constitutes a potent
neurite growth inhibitory protein (Chen et al., 2000; Prinjha
et al., 2000). Since the IgM as well as its recombinant F_{ab}
fragment effectively neutralize the inhibitory effect of CNS
myelin on axonal growth (Bandtlow et al., 1996; Broesamle
et al., 2000) we conclude that the exon 3 domain is involved
in the inhibitory activity of Nogo-A. This interpretation is
in agreement with observations from immunocytochemical
staining of live oligodendrocytes with an unrelated exon 3-
specific antibody (Huber and Schwab, 2000) and with func-
tional assays based on an exon 1–3 fragment that was expressed
as an IgG Fc fusion protein (Prinjha et al., 2000). However,
an inhibitory effect on axonal regeneration was recently
reported for a region called ‘Nogo-66’, which is found between
the two hydrophobic, presumably transmembrane, segments
within the C-terminal domain of Nogo-A (GrandPré et al.,
2000; Fournier et al., 2001). Therefore, regardless of the
current model of its transmembrane topology, it remains
possible that both the exon 3 region and the Nogo-66 region
inhibit axon growth, possibly via differing receptors.

The fact that the exon 3 moiety of Nogo-A was readily
secreted from E.coli, including efficient processing of the
bacterial signal peptide that was attached to its N-terminus,
and recovered as a soluble protein from the periplasmic space,
would be in agreement with the notion of its nature as an extracellular domain. We found that several of its eight Cys residues are important for proper folding, indicating that the large domain may be adapted to an oxidizing extracellular environment instead of the reducing cytosolic milieu. Even though the native full-length Nogo-A lacks a conventional signal peptide, which normally drives translocation of an N-terminal extracellular domain of a transmembrane protein across the membrane bilayer (von Heijne, 1996), peculiar post-translational mechanisms are known which may determine unusual topologies (Broome-Smith et al., 1994; Lu et al., 1998). Nevertheless, it remains to be seen from future experiments whether different orientations could exist for this atypical membrane protein. The engineered IN-1 F_{ab} fragment with its improved affinity should provide a helpful reagent in this respect.

The thorough characterization of the molecular interaction between the recombinant IN-1 F_{ab} fragment, as well as its improved affinity, on the other. Finally, the engineered plasticity effects with its moderate dissociation constant, enables site-directed ——

References

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IN-1 F_{ab} fragments with improved affinity for Nogo-A