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(54) Title: FUSION PROTEINS FOR THE TREATMENT OF CNS

(57) Abstract: This disclosure relates to compositions capable of use in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS), and in particular, compositions including proteoglycan degrading molecules and compositions capable of blocking and/or over coming the activity of neuronal growth inhibitory molecules, as well as fusion proteins which includes a proteoglycan degrading domain and a domain capable of blocking and or over coming the activity of neuronal growth inhibitory molecules.

FUSION PROTEINS FOR TREATMENT OF CNS

BACKGROUND AND SUMMARY

[0002] Spinal cord injury (SCI) inflicts trauma to the cells and tissues of the central nervous system (CNS) and causes a severe and debilitating condition in the individual. Following SCI, limited regeneration of injured neurons results in permanent disability characterized by some loss of sensation, paralysis and autonomic dysfunction. One reason that neurons fail to regenerate is their inability to traverse the glial scar that develops following SCI. This glial scar contains extracellular matrix molecules including chondroitin sulfate proteoglycans (CSPGs). In vitro studies show that neurons fail to extend processes over CSPG

coated surfaces, while in vivo data correlate failure of regeneration with areas of CSPG expression. Within the adult central nervous system (CNS) myelin there are also several identified axon growth inhibitor compounds like myelin associated glycoprotein (MAG), OMgp, and Reticulon 4 or Nogo that have been shown to be inhibitory for the growth of neurons.

[0003] The proteoglycan degrading enzyme chondroitinase ABC type I (**SEQ ID NO: X**) has been used to enhance neuronal growth in a dorsal column lesion model of spinal cord injury. It has also been reported that treating a spinal chord injury with NOGO receptor antagonist promotes a certain limited degree of neuronal regeneration. It has further been reported that creating a NOGO knock out mouse resulted in certain limited and inconsistent degrees of neuronal regeneration following dorsal hemisection of the spinal cord.

[0004] Experimental treatments for injury to the CNS have utilized the application of chondroitinase to the extracellular space, however the enzyme digests CSPG in the extracellular matrix and not intracellular stores. Furthermore, diffusion of chondroitinase within the parenchyma (the essential and distinctive tissue of an organ or an abnormal growth as distinguished from its supportive framework) or between anatomical compartments is limited. The limited access of drugs, imaging agents, and anesthetics, etc. to target cells and/or tissues of the Central Nervous System (CNS) may reduce the usefulness or effectiveness of any of these substances.

[0005] The delivery of therapeutic and diagnostic molecules into cells and tissues is, in part, dependent upon extracellular matrices as well as carbohydrates and proteins linked to cell membranes. The extracellular matrix is composed in part of proteoglycans, among them are the chondroitin sulfated proteoglycans (CSPGs). CSPGs are a family of proteoglycans composed of a core protein and covalently linked sulfated glycosaminoglycans. Each proteoglycan is

determined by the glycosaminoglycan side chains. For CSPGs these side chains are made up of approximately 40 to 100 sulfated disaccharides composed of chondroitin 4, 6 and dermatan sulfates. The protein component of the CSPG is ribosomally synthesized and the glycosylation occurs in the endoplasmic reticulum and Golgi apparatus. The sugar chains are then sulfated at the 4 or 6 positions by several glycosaminoglycan sulfotransferases.

[0006] Transduction proteins may be used to transport polypeptides and polynucleotides cargo across anatomical barriers and into cells. For example, the TAT protein from the human immunodeficiency virus (HIV) contains a protein transduction domain (PTD) that is involved in the transduction of HIV into cells. The PTD contains an 11 amino acid domain (TAT Peptide) that is responsible for the activity of the PTD. The TAT Peptide can be linked to proteins and facilitate the transduction of the proteins into cells. The mechanism of transduction is independent of the molecular weight or chemical properties of the proteins that are linked to the TAT Peptide. In vivo studies show that if a fusion protein consisting of the TAT Peptide linked to the 120 kd enzyme, beta-galactosidase (β -Gal), is injected into mice, then a robust delivery of β -Gal into a wide variety of cells is observed. When linked to the TAT transduction peptide, β -Gal activity was observed in the brain; without the TAT Peptide, β -Gal was not observed in the brain. Transport across the blood brain barrier is normally restricted to certain hydrophobic small molecules and particular low molecular weight lipophilic peptides. Transport of proteins as large as β -Gal into the brain is usually not possible without substantial disruption of the blood brain barrier, but the TAT Peptide facilitates transport while leaving the blood brain barrier intact.

[0007] Chimeric proteins, also called fusion proteins, are hybrid proteins which combine at least parts of two or more precursor proteins or polypeptides. Chimeric proteins may be

produced by recombinant technology, i.e. by fusing at least a part of the coding sequence of one gene to at least a part of the coding sequence of another gene. The fused gene may then be used to transform a suitable organism which then expresses the fusion protein.

[0008] Tat Peptide Complexes Frankel et al. (U.S. Pat. Nos. 5,804,604; 5,747,641; 5,674,980; 5,670,617; 5,652,122) discloses the use of Tat peptides to transport covalently linked biologically active cargo molecules into the cytoplasm and nuclei of cells. Frankel only discloses covalently linked cargo moieties that are (therapeutic, diagnostic or prophylactic), and does not teach or suggest the attachment of molecules that facilitate diffusion, plasticity, neurite growth, and axon regeneration. These molecules can include but are not limited to molecules that over come neurite out growth inhibition, or promote nerve growth such as soluble NOGO antagonists like NgR₂₇₋₃₁₁, neural cell adhesion molecules like L1, neurotrophic factors, growth factors, phosphodiesterase inhibitors, and inhibitors of MAG or MOG. Additionally, deletion mutants may be combined with other compounds that promote remyelination such as neuregulins (GGF2) and antibodies that promote remyelination. or proteoglycan degrading molecules to Tat peptides.

[0009] Regeneration following SCI is limited because of a variety of obstacles that include the deposition of CSPG at the glial scar, demyelination of axons, lack of trophic support and lack of axonal guidance. A single therapy directed against one aspect of SCI may not be as effective as a combinatorial approach. Fusion proteins with chondroitinase will allow combinatorial therapy with a single protein. Fusion partners for chondroitinase that will be constructed in this proposal were chosen from among proteins that have evidence for efficacy in SCI.

[0010] The use of a molecule that has the ability to both degrade extracellular matrix glycoproteins and to block or overcome the inhibitory nature of myelin components may be used to improve the ability of damaged neurons to grow or regenerate compared with either treatment alone. The proteoglycan degrading molecules may also be used advantageously to provide a method of facilitating access and diffusion of substances into cells or tissues through the use of at least one enzyme capable of cleaving proteoglycans and preferably degrading chondroitin sulfate proteoglycans (CSPG).

[0011] Embodiments of the present invention include compositions that comprise polypeptides which cleave proteoglycans, polypeptides that block and/or over come the activity of neuronal growth inhibitory molecules, or a combination of these. The compositions containing the proteoglycan degrading molecule or neuronal growth inhibitory molecules may also include molecules for transduction of the polypeptides across cell membranes and the blood brain barrier. The compositions may be used in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS). The compositions can be used in the regeneration of damaged neurological tissue and facilitate the diffusion and transport of therapeutic molecules capable of blocking and/or over coming the activity of neuronal growth inhibitory molecules into damaged or diseased tissue.

[0012] Embodiments of the present invention include compositions and methods for their use to facilitate delivery and diffusion of therapeutics or diagnostic agents, and preferably agents that promote regeneration of nerves and axons, into cells or tissues. Preferably the composition includes the use of an enzyme capable of cleaving chondroitin sulfate proteoglycans (CSPG) to increase the diffusion of these agents into cells or tissues of the central nervous system.

[0013] Compositions of the present invention may include chimeric or fusion proteins capable of systemic use in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS), and in particular, fusion proteins capable of crossing the blood brain barrier. The fusion protein may include a polypeptide transduction domain, a polypeptide domain capable of degrading a proteoglycan, preferably a domain cleaving chondroitin sulfate proteoglycan (CSPG), a polypeptide domain that blocks and or over comes the activity of neuronal growth inhibitory molecules, or any combination of these polypeptide domains that may be used in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS) . The various polypeptide domains may be linked or chemically bonded together by polypeptide linkers.

[0014] Compositions of the present invention include polynucleotides which encode for the chimeric or fusion proteins capable of systemic use in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS), and in particular, they encode for fusion proteins capable of crossing the blood brain barrier. The polynucleotides which encode for these chimeric or fusion proteins may include a polynucleotide domain encoding for a polypeptide transduction domain, a polynucleotide domain encoding for a polypeptide domain capable of degrading a proteoglycan, preferably cleaving chondroitin sulfate proteoglycan (CSPG), a polynucleotide domain encoding for a polypeptide domain that blocks and or over comes the activity of neuronal growth inhibitory molecules, or any combination of these domains that may be used in the treatment of spinal cord injuries and related disorders of the central nervous system (CNS) . The polynucleotide also includes one or more polynucleotide domains that encode for polypeptides that link the domains of the polypeptide together to form the fusion protein.

[0015] One embodiment of the present invention is a composition and a method for its use that facilitates the access and distribution of therapeutic and diagnostic agent in the composition into cells, through membranes or into tissues by the use of composition that includes at least one enzyme capable of cleaving proteoglycans, preferably the composition includes a fusion protein having an enzyme capable of cleaving CSPG's. The molecules or agents in the composition may include one or more of Growth factors including, Brain Derived Neurotrophic Factor, Insulin-like Growth Factor, Fibroblast Growth Factor, Ciliary Neurotrophic Factor, Glial Derived Neurotrophic Factor, Transforming Growth Factor, Glial Growth Factor 2, L1, GM1, Vascular Endothelial Growth Factor, Nerve Growth Factor, Immunophilins. Molecules in the composition can include fluorescent or contrast agents for imaging. The agents may include cells for transplant – stem cells, neurons, others, cells as delivery agents, chemotherapeutic agents, antibiotics, antibody therapies, Nogo receptor antagonists, other chondroitinase enzymes. The composition may include a transduction domain, an enzyme capable of cleaving proteoglycans, or both. Preferably the composition includes a fusion protein having a transduction domain, an enzyme domain capable of cleaving proteoglycans, or both. The fusion protein can facilitate the transport or modifies transport of such agents into cells, tissues, and/or otherwise inaccessible locations; and/or to enhance penetration rates, distance of penetration; or provide more even concentration distribution. Preferably the modified transport occurs through the use of at least one enzyme capable of cleaving CSPGs. The compositions can be used for treating a CNS injury, preferably the composition is used in the treatment of neuronal damage from a contusion injury.

[0016] Embodiments of the present invention include chimeric proteins of a proteoglycan degrading domain linked to a polypeptide that blocks the action of neuronal growth inhibitors

such as but not limited to a Nogo-receptor antagonist (NgR₂₇₋₃₁₁) domain or variant linked to a chondroitinase like chondroitinase ABC I or a variant of chondroitinase having one or more N terminal amino acids deleted. The compound may include chimeric proteins of a proteoglycan degrading domain linked to a polypeptide that is a neural cell adhesion promoter such as an L1 neural cell adhesion promoter domain or variant linked to chondroitinase ABC I or a variant of chondroitinase having one or more N terminal amino acids deleted. The chimeric proteins may include chimeric proteins of a proteoglycan degrading domain linked to a polypeptide that is a glial cell stimulator, such as but not limited to a GGF2 glial cell stimulator or variant linked to chondroitinase ABCI or a variant of chondroitinase having one or more N terminal amino acids deleted.

[0017] An *E.Coli* recombinant expression system and purification process can be used to produce essentially pure and catalytically active chondroitinase ABCI. These methods may be modified for producing chimeras of proteoglycan degrading molecules and other agents.

[0018] The chimera may be assayed for chondroitinase enzymatic activity and the specific biological activity of each fusion partner. Methods to measure the activities of the chimera may be modified for those used to measure chondroitinase activity including a spectrophotometric assay, zymography, an HPLC assay to detect CSPG disaccharide digestion products and an in vitro neurite outgrowth assay. A neuron growth cone collapse assay can be used to evaluate NOGO receptor antagonists and a neurite outgrowth assay can be used measure LI activity. GGF2 activity may be measured using a Schwann cell proliferation assay.

[0019] The compositions and method of the present invention can be used for the treatment of spinal cord injuries and in the promotion of regeneration of axons. The compositions of the present invention can also be used to promote plasticity, regrowth, repair,

and/or regeneration of dysfunctional neurons in the CNS that have been damaged as a result of disease, such as degenerative diseases including Alzheimer's and Parkinson's disease.

Advantageously, the use of proteoglycan degrading polypeptides or membrane transducing polypeptides in the compositions of the present invention also promote diffusion and access of damage or diseased tissue to other therapeutic agents promoting the regeneration of neurons.

DESCRIPTION OF THE DRAWINGS

[0020] The file of this patent contains at least one drawing/photograph executed in color. Copies of this patent with color drawing(s)/photograph(s) will be provided by the Office upon request and payment of the necessary fee.

[0021] In part, other aspects, features, benefits and advantages of the embodiments of the present invention will be apparent with regard to the following description, appended claims and accompanying drawings where:

[0022] FIG. 1 is an illustrative example of a TAT-chondroitinase ABCI construct; the DNA sequence for the entire gene fragment fused with the Tat sequence followed by the 5'-glycine linker; the sense and antisense oligonucleotides having the sequences as 5'-tatgtatggtc gtaaaaagcgctc gtcaacgtcgtcgtgg tggtggtggta-3' and 5'-tatgaccaccaccaccacgac gacgttgacgac gctttt acgaccataca-3' were annealed and ligated at the NdeI site which is at the 5' end of ABCI gene cloned in pET15b (Novagen) at the NdeI and BamHI sites. The Tat₄₈₋₅₇ peptide sequence (GRKKRRQRR) is joined with the ABCI sequence by a pentaglycine linker.

[0023] FIG. 2 are Brain section images; (I) illustrating lobes from adult rat which were incubated in beta-galactosidase alone (B&D), or with the addition of Chondroitinase ABCI (A, 0.5 U/ml or C, 0.005 U/ml); (II) Eosin Y penetration through the cortex of a Chondroitinase treated brain hemisphere and control showing approximately the same penetration, Eosin Y is a

is a zwitterionic, having an overall negative charge at the low pH it was used at, and it is 692 kDa; (III) A saturate solution of Congo Red demonstrates greater penetration through the cortex of a Chondroitinase treated brain hemisphere as compared to untreated brain, Congo red is a negatively charged dye of 697 kDa;

[0024] FIG. 3 (A) is a diagram representing the full length GGF protein (GGF2-M₁-E₄₂₂) and the three GGF2 fragments containing the Ig and EFG domains alone and in combination; (B) a schematic presentation of chimeric proteins of a proteoglycan degrading molecule like chondroitinase ABCI and a neuregulin 1 gene isoform GGF2;

[0025] FIG. 4 illustrate the structure of (A) an NgR₂₇₋₃₁₁- N terminal chondroitinase ABCI chimeric protein with and without a peptide spacer or linking group; (B) an NgR₂₇₋₃₁₁- C terminal chondroitinase ABCI chimeric protein with and without a peptide spacer or linking group; (C) an extracellular domain L1 N-terminal chondroitinase ABCI chimeric protein with and without a spacer or linking peptide; (D) an extracellular domain L1 C-terminal chondroitinase ABCI chimeric protein with and without a spacer or linking peptide.

[0026] FIG. 5 (A) BBB scores from spinal cord injured animals treated with chondroitinase (Chondroitinase ABC I), penicillinase or artificial cerebrospinal fluid (aCSF); (B) Parasagittal sections of spinal cords from injured animals treated with chondroitinase (left column) or penicillinase (right column). Sections were cut at 30 microns. Each set of images contains four parasagittal sections realigned with the rostral cord to the left and the caudal cord to the right. The most central section of each set has been removed and replaced below to aid visualization. Pairs of images are Weil stained, anti-GFAP and an amino cupric stain of neuronal degeneration (top to bottom). (C) and (D) Distribution of individual BBB scores according to

treatment group. Scores are BBB scores at ten weeks post surgery. Group averages are shown below the data points.

DETAILED DESCRIPTION

[0027] Before the present compositions and methods are described, it is to be understood that this invention is not limited to the particular molecules, compositions, methodologies or protocols described, as these may vary. It is also to be understood that the terminology used in the description is for the purpose of describing the particular versions or embodiments only, and is not intended to limit the scope of the present invention which will be limited only by the appended claims.

[0028] It must also be noted that as used herein and in the appended claims, the singular forms "a", "an", and "the" include plural reference unless the context clearly dictates otherwise. Thus, for example, reference to a "cell" is a reference to one or more cells and equivalents thereof known to those skilled in the art, and so forth. Unless defined otherwise, all technical and scientific terms used herein have the same meanings as commonly understood by one of ordinary skill in the art. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of embodiments of the present invention, the preferred methods, devices, and materials are now described. All publications mentioned herein are incorporated by reference. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure by virtue of prior invention.

[0029] "Optional" or "optionally" means that the subsequently described event or circumstance may or may not occur, and that the description includes instances where the event occurs and instances where it does not.

[0030] Suitable enzymes that cleave CSPGs include chondroitinase ABC type I, chondroitinase ABC Type II, chondroitinase AC and Chondroitinase B or mammalian enzymes with chondroitinase-like activity such as hyaluronidase 1, hyaluronidase 2, hyaluronidase 3, hyaluronidase 4, cathepsins, ADAMTs and PH-20 or mixtures thereof.

[0031] CSPGs are a family of proteoglycans composed of a core protein and covalently linked sulfated glycosaminoglycans. The polysaccharide chains are cleaved by several enzymes, including a family of chondroitinases. To date, this enzyme family contains at least four members, Chondroitinase ABC I, ABC II, AC and B. Chondroitinase ABC I is an exo-lyase which cleaves both chondroitin and dermatan sulfates. It has been used extensively in the study of in vitro neuronal regeneration over CSPG-laden substrates and more recently in vivo studies following CNS injury.

[0032] Proteins and polypeptide that may be used in the compositions and fusion proteins of the present invention are those which promote plasticity as well as the regeneration of injured or diseased neurons and axons. These regenerating proteins and polypeptides may include cell adhesion proteins, those which stimulate glial cells, and polypeptide which block the inhibitory effect of proteins that act as axon growth inhibitors.

[0033] Plasticity of the nervous system refers to any type of functional reorganization. This reorganization occurs with development, learning and memory and brain repair. The structural changes that occur with plasticity may include synapse formation, synapse removal, neurite sprouting and may even include strengthening or weakening existing synapses. Regeneration is generally differentiated from plasticity by the long range growth of axons in disrupted tracts that is characteristic of regeneration.

[0034] Proteins and polypeptides are capable of blocking the activity of neuronal growth inhibitory molecules may include peptides and polypeptides that block the inhibitory properties of peptide such as but not limited Nogo, MAG, OMgp. Suitable compositions that overcome the activity of neuronal growth inhibitory molecules include but are not limited to Protein Kinase C family inhibitors, Rho Kinase family inhibitors and agents, such as phosphodiesterase inhibitors, that increase intracellular cyclic AMP, and L1. The (NgR₂₇₋₃₁₁) peptide has been shown to inhibit binding of Nogo66, OMgp, MAG, and MOG to membrane-bound NogoR and overcome the inhibitory effects of Nogo on the regeneration of nerve processes.

[0035] Proteins and polypeptides that affect cell adhesion or stimulate cells may include but are not limited to poly peptides such as L1 and GGF2. L1 is a neural cell adhesion protein that is a potent stimulator of neurite outgrowth in vitro for which has been found that treatment of acute SCI in rodents using a soluble form of L1 leads to an increase in the recovery of neurological function. GGF2 stimulates glial cells and has been shown to improve clinical outcome measures in a murine model of experimental allergic encephalomyelitis (EAE) likely as a result of the stimulation of oligodendrocytes to promote remyelination.

[0036] Cell membrane-permeant peptide sequences useful in practicing the present invention include, but are not limited to, RQARRNRRRRWRERQR-51 (HIV-1 Rev protein basic motif; (SEQ ID NO:51)); MPKTRRRPQRSQRKPPTP-119 (HTLV-1 Rex protein basic motif; SEQ ID NO:52)) (Kubota et al. 1989); the third helix of the homeodomain of Antennapedia (Derossi, et al., J. Biol. Chem. 271:18188-93, 1996) (43-RQILIWFQNRRMKWLL-58 (SEQ ID NO:53)); a peptide derivable from the heavy chain variable region of an anti-DNA monoclonal antibody (Avrameas, et al., Proc. Natl. Acad. Sci. 95:5601-06, 1998) (VAYISRGGVSTYYSDTVKGRFTRQKYNKRA (SEQ ID NO:54)); and the

Herpes simplex virus VP22 protein (Elliot and O'Hare, Cell, 88:223-33, 1997) (1-MTSRRSVKSGPREVPRDEYEDLYYTPSSGMASPDSPPDTSRRGALQTRSRQRGEVRFVQYDESDYALYGGSSSEDDEHPEVPRTRRPVSGAVLSGPGPARAPPPPAGSGGAGRTPTTAPRAPRTQRVATKAPAAPAAETTRGRKSAQPESAALPDAPASRAPTVQLWQMSRPRTDEDLNELLGITHRVTVCEGKNLLQRANELVNPDVVQDVDAATATRGRSAASRPTERPRAPARSASRPRRPVE-246 (SEQ ID NO:55)). In a preferred embodiment, the basic peptide is derivable from the human immunodeficiency virus type 1 (HIV-1) Tat protein (Fawell et al., Proc. Natl. Acad. Sci., 91:664-68, 1994). In particular, the Tat peptide can comprise any sequential residues of the Tat protein basic peptide motif 37-72 (Vives et al. 1997) (37-CFITKALGISYGRKKRRQRRRPPQGSQTHQVSLSKQ-72 (SEQ ID NO:56)). The minimum number of amino acid residues can be in the range of from about three to about six, preferably from about three to about five, and most preferably about four, i.e., the minimal requirement for one alpha helical turn. A preferred embodiment comprises Tat protein residues 48-57 (GRKKRRQRRR) (SEQ ID NO:57).

[0037] Suitable PTD domains include those derived from the TAT protein. Tat Proteins and Peptides Tat is an 86-amino acid protein involved in the replication of human immunodeficiency virus type 1 (HIV-1). The HIV-1 Tat transactivation protein is efficiently taken up by cells (Mann and Frankel 1991; Vives et al. 1994), and low concentrations (nM) are sufficient to transactivate a reporter gene expressed from the HIV-1 promoter (Mann and Frankel 1991). Exogenous Tat protein is able to translocate through the plasma membrane and reach the nucleus to transactivate the viral genome.

[0038] A region of the Tat protein centered on a cluster of basic amino acids is believed to be responsible for this translocation activity (Vives et al. 1997). Tat peptide-mediated cellular

uptake and nuclear translocation have been demonstrated in several systems (Vives, et al., J Biol Chem 272:16010-16017, 1997; Jones, Genes Dev 11:2593-2599, 1997). Chemically coupling a Tat-derived peptide (residues 37-72) to several proteins results in their internalization in several cell lines or tissues (Fawell, et al., Proc Natl Acad Sci USA 91:664-668, 1994; Anderson, et al., Biochem Biophys Res Commun 194:876-8884, 1993; Fahraeus, et al., Curr Biol 6:84-91, 1996; Nagahara, et al., Nat Med 4:1449-1452, 1998). A synthetic peptide consisting of the Tat basic amino acids 48-60 with a cysteine residue at the C-terminus coupled to fluorescein maleimide translocates to the cell nucleus as determined by fluorescence microscopy (Vives et al. 1997). In addition, a fusion protein (Tat-NLS-.beta.-Gal) consisting of Tat amino acids 48-59 fused by their amino-terminus to .beta.-galactosidase amino acids 9-1023 translocates to the cell nucleus in an ATP-dependent, cytosolic factor-independent manner (Efthymiadis et al. 1998).

[0039] Chimeric proteins, also referred to in the art as fusion proteins, are hybrid proteins which combine at least parts of two or more precursor proteins or peptides. Chimeric proteins may be produced by recombinant technology, i.e. by fusing at least a part of the coding sequence of one gene to at least a part of the coding sequence of another gene. Where desirable, one or more genes for linker peptides may be fused to the coding sequence of genes for the other polypeptide domains in the fusion protein. The fused gene may then be used to transform a suitable organism such as but not limited to *E. coli* or CHO cells which then expresses the fusion protein.

[0040] Genes encoding either N or C terminal deletion mutants of polypeptide domains of the fusion proteins can be used in constructs for expression of the fusion proteins. Preferably the generated deletion mutants maintain their catalytic proteoglycan degrading activity, blocking activity, growth activity, or transduction activity. Generated deletion mutants of the

proteoglycan degrading molecules like chondroitinase ABCI enzyme where the mutant is missing a certain number of amino acids from the N and or C-terminal are those that retain some proteoglycan degrading activity. N-terminal deletions of chondroitinase like chondroitinase ABC I maintain a histidine-tag that is attached to the N-terminus. It is expected that a TAT-deletion mutant chondroitinase ABC I fusion DNA construct can be expressed without removal of the TAT polypeptide during expression. For example Fuse TAT peptide at the N-terminus of either ABCI-NΔ20 or ABCI-NΔ60 deletion mutant. Fragments of polypeptides, such as those shown in FIG. 3 for GGF2, may be used in the construction of chimeric fusion proteins.

[0041] Catalytically active deletion mutants of chondroitinase ABCI can be prepared for example but not limited to deleting 20, 40 and 60 amino acids respectively from the N-terminus of the mature ABCI protein. Deletion of 60 amino acids from the N terminal and 80 amino acids from the C-terminal end may also be used to make a deletion mutant of a proteoglycan degrading chondroitinase ABCI. These deletion mutants and those of other proteoglycan degrading molecules may be used for construction of N-terminal fusion chimeric protein. Detailed comparative biochemical studies can be done to determine the efficacy of mature chondroitinase ABCI versus various deletion mutant in compositions and fusion proteins with respect to the substrate specificity, substrate binding and tissue penetration.

[0042] A mutant of chondroitinase ABCI that has native protein structure, but lacks catalytic activity may be prepared as a null or a negative control for bioassays and SCI studies. Based on the crystal structure of chondroitinase ABCI a site-specific mutant designated H501a and Y508a to knock out catalytic activity in the putative active site can be prepared. Such mutants can be tested for inactivation of catalytic activity and SEC to compare to the wild-type

enzyme. If the null activity mutant is successfully created it will provide a negative control for the various fusion proteins for use in bioassays and ultimately in SCI animal studies.

[0043] An *E.Coli* expression system may be used to make chondroitinase using PET expression vectors (Novagen). The GGF2-chondroitinase ABCI fusion protein may be expressed in *E.coli*. Constructs for Tat-chondroitinase deletion mutant fusion proteins, Tat-GGF2 fusion proteins, or Tat-chondroitinase-GGF2 fusion proteins may be expressed from *E.coli*. Other fusion proteins can be expressed in CHO cell lines.

[0044] Table 1 illustrates various non-limiting components which may be used in compositions of the present invention as a mixture, and preferably as a fusion or chimeric molecule. The composition or chimeric molecule described may include one or more of the molecules in Table 1. In the case of chimeric molecules one or more linker segments, preferably polypeptides are used.

Table 1 Components of Compositions

Transduction molecules	Proteoglycan degrading molecules	Therapeutic, diagnostic, receptor antagonist molecules
Tat protein residues 48-57 HIV-1 Rev protein basic motif HIV-1 Rev protein basic motif Herpes simplex virus VP22 protein	Any agent that degrades extracellular matrix glycoproteins including: Chondroitinase ABC I, Chondroitinase ABC II, Chondroitinase AC, Chondroitinase B, Hyaluronidase 1, Hyaluronidase 2, Hyaluronidase 3, Hyaluronidase 4, PH20 deletion and or substitution mutants of the above listed molecules that maintains enzymatic activity.	Any peptide that blocks the inhibitory properties of Nogo, MAG, OMgp including: Nogo peptide 1 – 40* Any component of Nogo peptide 1-40 that maintains the ability to block the inhibitory properties of Nogo Other blocking peptides of Nogo Antibodies that recognize Nogo Blocking peptides of MAG Antibodies that recognize MAG Blocking peptides of OMgp Antibodies that recognize OMgp Antibodies that recognize the

		<p>Nogo-66 receptor Blocking peptides of the p75 receptor Antibodies that recognize the p75 neurotrophin receptor Peptides or antibodies that block other receptors of Nogo, MAG and/or OMgp</p> <p>Peptide that overcomes the inhibitory properties of Myelin, Nogo, MAG, OMgp including: Protein Kinase C family inhibitors* Rho Kinase family inhibitors Agents that increase intracellular cAMP concentration L1</p>
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*Nogo peptide 1-40: The human 40-aa NEP1-40 aa (Acetyl-RIY KGV IQA IQK SDE GHP FRA YLE SEV AIS EEL VQK YSN S-amide corresponding to 1-40 aa of Nogo-66

[0045] A schematic illustration of non-limiting versions of chimeric fusion proteins of the present invention are illustrated in FIG. 4A and FIG. 4B.

[0046] A peptide component from any column in Table 1 could be linked by an oligopeptide linker well known in the art such as a glycine rich peptide, for example Gly-Gly-Gly-Gly, or linkers prepared including one of the naturally occurring amino acids as well as substituted or beta or gamma amino acids like 4-aminobutyric acid or 6-aminocaproic acid can be used. Other linkers including but not limited to alkyl diamines, amino, or alkyl diols may also be used. Preferably the Transduction component of the fusion protein is in a terminal position in the chimeric protein. Other examples of common linkers may include but would not be limited to Gly-Gly-Ala-Gly-Gly, Gly/Ser rich linkers (for example Gly₄Ser₃), or Gly/Ala rich linkers.

Additionally, linkers may be of any length and design to promote or restrict the mobility of components in the fusion protein.

[0047] The incorporation of non-natural amino acids, including synthetic non-native amino acids, substituted amino acids, or one or more D-amino acids into the peptides (or other components of the complexes) of the present invention (subsequently referred to herein as "D-peptides") is advantageous in a number of different ways. D-amino acid-containing peptides exhibit increased stability in vitro or in vivo compared to L-amino acid-containing counterparts. Thus, the construction of peptides incorporating D-amino acids can be particularly useful when greater intracellular stability is desired or required. More specifically, D-peptides are resistant to endogenous peptidases and proteases, thereby providing better oral transepithelial and transdermal delivery of linked drugs and conjugates, improved bioavailability of membrane-permeant complexes, and prolonged intravascular and interstitial lifetimes when such properties are desirable. The use of D-peptides can also enhance transdermal and oral transepithelial delivery of linked drugs and other cargo molecules.

[0048] In a spinal cord injury, the axons of ascending sensory and descending motor neurons are disrupted, that can result in the loss of sensation and paralysis. These axons fail to regenerate successfully leading to permanent disability. A scar envelopes the site of the injury which is believed to wall off the area of fragile tissue, stabilize the blood brain barrier, and prevent an overwhelming cascade of uncontrolled tissue damage. This scar is composed of hypertrophic glial cells and an extracellular matrix (ECM). Chondroitin sulfate proteoglycans (CSPGs) are one important component of the scar. They are expressed by glial cells and deposited in the ECM in regions of blood brain barrier breakdown. *In vitro* evidence demonstrates that these CSPGs are potently inhibitory for the growth of axons and without

wishing to be bound by theory, are believed to contribute to the failure of the spinal cord axons to regenerate and reform functional synapses. *In vivo* studies have demonstrated that regenerating axons are able to grow into and even beyond the scar.

[0049] CSPGs and white matter components are generally accepted as molecular barriers that neurons must overcome in order to regenerate and reestablish functional connections after injury. Transplanted adult sensory neurons placed distal to a forming scar can regenerate robustly even along degenerating white matter pathways, however, regeneration ceases abruptly as axons enter the proteoglycan containing glial scar. Treatment of CNS white matter pathways with chondroitinase enhances the ability of neurons to grow in these substrates.

[0050] Central nervous system tissues are tightly compacted with cells and have limited extracellular space. The proteins and carbohydrates of the extracellular matrix provide charge and osmotic forces as well as specific and non-specific binding sites which may prevent the penetration of therapeutic agents. The enzymatic cleavage of these matrix and cellular components may later or facilitate the access of compounds or cells through tissues. A proteoglycan degrading molecule like Chondroitinase ABC I that is an enzyme that digests chondroitin sulfate proteoglycans can be used to promote diffusion of therapeutic molecules into the CNS. Tat peptides to transport covalently linked biologically active cargo molecules into the cytoplasm and nuclei of cells. In the case of a fusion protein having a protein transduction polypeptide domain like the HIV tat protein, therapeutic molecules for axon regeneration may be delivered across the blood brain barrier.

[0051] Treatment of SCI model injuries, preferably contusion model injury, can be used to determine the degree of regeneration and functional recovery achieved by compositions and method of the present invention. The degree of functional recovery can be demonstrated by

improved corticospinal tract conduction, improved tape removal, beam walking, grid walking and paw placement following chondroitinase treatment of a dorsal column lesion. Motor skill improvement as well as autonomic function: bowel, bladder, sensor and sexual function may also be used as measures of function improvement and related to molecular structure and components in the compositions of the present invention.

[0052] In addition to the ability of chondroitinase to enhance regeneration, chondroitinase digests components of the perineuronal network (PNN). The density of the PNN is variable within the CNS and is particularly dense in the somatosensory, auditory, visual cortices and the hippocampus. PNN has also been demonstrated to be dense around spinal motor neurons. The inventors have discovered the dense PNN within the dorsal horn of the cord. Digestion of the PNN can enhance plasticity within the hippocampus and visual cortex. Plasticity within intact systems in incomplete SCI, especially in the region of the central pattern generators or in the reticular core, may support the function of damaged or destroyed systems. Promotion of plasticity in these systems may be one mechanism other than or in addition to regeneration by which chondroitinase can improve function following CNS injury. Furthermore, regeneration and plasticity may work in concert to affect recovery following injury; indeed, the corticospinal tract has been shown to be critical for modulation of spinal cord plasticity.

[0053] Recovery of neurological function following contusion injury in the CNS or a disease state may be promoted by administering the fusion proteins or mixtures including one or more of the components in Table 1, to cells, a tissue, or a subject having damaged or diseased neurons whether the injury or disease is immediate or long-standing.

[0054] The fusion proteins herein are administered in an amount effective to degrade CSPGs and thereby promote the recovery of neurological function. Once the proteins or

polypeptides in the compositions have been purified to the extent desired, they may be suspended or diluted in an appropriate physiological carrier or excipient for SCI treatment. In models of SCI, effective intrathecal doses in rats have been about 0.06 units on alternate days for 14 days. A dose for a 70 kilogram human may be about 17 Units. At about 100 Units / milligram, this would equal about 170 micrograms. Doses of up to 20 Units appear safe in the rat. Compositions including a proteoglycan degrading molecule in a mixture or as part of a fusion protein diluted in a carrier or pharmaceutically acceptable excipient can be injected, generally at concentrations in the range of 1 µg to 500 mg/kg of host. Administering the agent can be by bolus injection, intravenous delivery, continuous infusion, sustained release from implants, or sustained release pharmaceuticals. Administration may be by injection, such as intramuscularly, peritoneally, subcutaneously, intravenously. Oral administration may include tablets or capsules, preferably the oral dosage is a sustained release formulation for once or twice daily administration. Percutaneous administration can be once per day, and is preferably less than once per day administration . Administration to the human patient or other mammalian subject may be continued until a measurable improvement in autonomic or motor function in the patient is achieved.

[0055] The chondroitinase PTD fusion proteins can be administered with a suitable pharmaceutical carrier. The administration of the compositions of the present invention as mixtures or chimeric proteins can be topical, local or systemic. The chimeric fusion proteins may also be secreted by genetically cells, preferably a chimeric fusion protein having a proteoglycan degrading portion like chondroitinase and a transduction polypeptide potion like TAT can be secreted by genetically modified cells that are implanted, either free or in a capsule, at or near the site of CNS injury.

[0056] Once the compositions either as mixtures or fusion proteins are administered, degradation of CSPGs removes the inhibitory molecules that block neurite outgrowth, and allow the regeneration of neurites into the affected area. For example, the chondroitinase AC and chondroitinase B degrade CS and DS, respectively, resulting in unsaturated sulfated disaccharides. Chondroitinase AC cleaves CS at 1, 4 glycosidic linkages between N-acetylgalactosamine and glucuronic acid in the polysaccharide backbone of CS. Cleavage occurs through beta-elimination in a random endolytic action pattern. Chondroitinase B cleaves the 1, 4 galactosamine iduronic acid linkage in the polysaccharide backbone of DS. The cleavage of both CS and DS occurs through a beta-elimination process which differentiates these enzymatic mechanisms from mammalian GAG degrading enzymes. Chondroitinase ABCI, chondroitinase ABCII, are exo and endo lyases that cleave both CS and DS. The removal of CS and DS from the glial scar permits the regeneration of neurite outgrowths into the injured area.

[0057] Mixtures of any of these fusion polypeptides may be used to provide a therapeutic treatment for CNS injuries and disorders which may include but not limited to contusion injury, traumatic brain injury, stroke, multiple sclerosis, brachial plexus injury, ambliopia, spinal cord injuries. Spinal cord injuries includes disease and traumatic injuries, such as the crushing of neurons brought about by an auto accident, fall, contusion, or bullet wound, as well as other injuries. Practice of the present methods can confer clinical benefits to the treated mammal, providing clinically relevant improvements in at least one of the subject's motor coordination functions and sensory perception. Clinically relevant improvements can range from a detectable improvement to a complete restoration of an impaired or lost function of the CNS.

[0058] The regeneration of the nerve cells in to the affected CNS area allows the return of motor and sensory function. Clinically relevant improvement will range from a detectable

improvement to a complete restoration of an impaired or lost nervous function, varying with the individual patients and injuries.

[0059] A variant of a protein or fragments thereof refer to a molecule substantially similar to either the entire protein or a fragment, which possesses biological activity that is substantially similar to a biological activity of the complement protein or fragments. A molecule is substantially similar to another molecule if both molecules have substantially similar structures or if both molecules possess a similar biological activity.

[0060] Variants of complement protein or fragments thereof are produced by chemical or recombinant means. Variants of the polynucleotides constructed to express fusion proteins may also be made. The variants may include, for example, deletions from, or insertions or substitutions of, amino acid residues within the amino acid sequence, or deletion, substitution, or insertion of nucleic acids from a sequence encoding for a particular fusion protein or polypeptide domain in the fusion protein . For example, in some cases the removal of one or more amino acid residues from a chondroitinase polypeptide can be made without significant change in its CSPG degradation activity. Substantial changes in functional properties like proteoglycan degradation or blocking activity against axon growth inhibitors are made by selecting substitutions that are less conservative, ie. that differ more significantly in their effect on maintaining the structure, charge or hydrophobicity of the peptide backbone in the area of the substitution.

[0061] Most deletions, insertions, and substitutions are not expected to produce radical changes in the characteristics of the protein molecule; however, when it is difficult to predict the exact effect of the substitution, deletion, or insertion in advance of doing so, one skilled in the art will appreciate that the effect will be evaluated by routine screening assays. For example, a

change in the axon regeneration character of the polypeptide molecule, through more or less proteoglycan degradation can be measured with functional recovery tests as well as HPLC assays to detect CSPG disaccharide digestion products.

[0062] The present invention relates to the use of a proteoglycan degrading molecule, an HIV tat protein, or a tat-derived polypeptide, or a combination of these as a mixture or fusion protein to deliver a molecule of interest into the CNS or a site in the CNS where neuronal tissue damage has occurred by disease or trauma. In particular the damage site in the CNS is where scarring has occurred as a result of a contusion injury. The molecule of interest, optionally referred to as an agent or cargo molecule can be a therapeutic molecule which promotesplasticity, axon growth, a diagnostic molecule, or a proteoglycan degrading molecule. In the case of a fusion protein having a protein transduction polypeptide domain like the HIV tat protein, therapeutic molecules for axon regeneration may be delivered across the blood brain barrier or proteoglycan degrading molecule can be delivered into cells to degrade cellular stores of proteoclycans and promote axon regeneration.

[0063] Transport polypeptides of this invention may be advantageously attached to cargo molecules by chemical cross-linking or by genetic fusion. A unique terminal cysteine residue is a preferred means of chemical cross-linking. According to some preferred embodiments of this invention, the carboxy terminus of the transport moiety is genetically fused to the amino terminus of the cargo moiety. Embodiment of the present invention consists of an amino-terminal methionine followed by tat residues 47-58, followed by a chondroitinase polypeptide.

[0064] It will be appreciated that the entire 86 amino acids which make up the tat protein may not be required for the uptake activity of tat. For example, a protein fragment or a peptide which has fewer than the 86 amino acids, **but which exhibits uptake into cells and or can**

cross the blood brain barrier, can be used (a functionally effective fragment or portion of tat).

For example, tat protein containing residues 1-72 can be sufficient for uptake activity and tat residues 1-67 can mediate the entry of a heterologous protein into cells. Synthetic peptide containing tat residues 1-58 can have uptake activity.

[0065] The tat peptide can be a single (i.e., continuous) amino acid sequence present in tat protein or it can be two or more amino acid sequences which are present in tat protein, but in the naturally-occurring protein are separated by other amino acid sequences. As used herein, tat protein includes a naturally-occurring amino acid sequence which is the same as that of naturally-occurring tat protein, its functional equivalent or functionally equivalent fragments thereof (peptides). Such functional equivalents or functionally equivalent fragments can possess uptake activity into the cell or across the blood brain barrier that is substantially similar to that of naturally-occurring tat protein. Tat protein can be obtained from naturally-occurring sources or can be produced using genetic engineering techniques or chemical synthesis.

[0066] The amino acid sequence of naturally-occurring HIV tat protein can be modified, by addition, deletion and/or substitution of at least one amino acid present in the naturally-occurring tat protein, to produce modified tat protein (also referred to herein as tat protein or polypeptide). Modified tat protein or tat peptide analogs with increased stability can thus be produced using known techniques. Therefore, tat proteins or peptides may have amino acid sequences which are substantially similar, although not identical, to that of naturally-occurring tat protein or portions thereof. In addition, cholesterol or other lipid derivatives can be added to tat protein to produce a modified tat having increased membrane solubility.

[0067] Naturally-occurring HIV-1 tat protein has a region (amino acids 22-37) wherein 7 out of 16 amino acids are cysteine. Those cysteine residues are capable of forming disulfide

bonds with each other, with cysteine residues in the cysteine-rich region of other tat protein molecules and with cysteine residues in a cargo protein or the cargo moiety of a conjugate. Such disulfide bond formation can cause loss of the cargo's biological activity. Furthermore, even if there is no potential for disulfide bonding to the cargo moiety (for example, when the cargo protein has no cysteine residues), disulfide bond formation between transport polypeptides can lead to aggregation and insolubility of the transport polypeptide, the transport polypeptide-cargo conjugate, or both. The tat cysteine-rich region may be deleted to avoid disulfide bond formation and prevent aggregation and insolubility of the transport polypeptide, the transport polypeptide-cargo conjugate, or both.

[0068] Chondroitinase is also able to promote plasticity in regions of the CNS with significantly dense PNN, including cortex, tectum, hippocampus and spinal cord. It is reasonable that some combination of effects, including regeneration, sprouting and plasticity are responsible for the improvement in function following SCI with chondroitinase treatment or treatment with fusion molecules including chondroitinase or other proteoglycan degrading molecule.

[0069] NbR₂₇₋₃₁₁: Nogo is a high molecular weight myelin component that inhibits neurite outgrowth. The amino terminal region (Nogo66) is the part of the molecule that is specifically associated with the inhibition of neurite outgrowth. Expression cloning methods revealed the receptor for Nogo66 (NgR) is a GPI anchored glycoprotein expressed mainly on neurons. NgR interacts not only with Nogo, but also with other myelin associated inhibitors such as MAG and MOG. Due to its central role in the inhibitory properties of myelin on neurons, NgR has been the target for approaches to antagonize its interactions with its ligands. The soluble segment of NgR interacts with Nogo66, MAG and MOG. The region spans residues

27-311 and this NgR fragment can therefore act as a decoy receptor that will interfere with myelin associated inhibition of neurons. If NgR₂₇₋₃₁₁ is linked to proteoglycan degrading molecule like chondroitinase ABCI to form a chimeric fusion protein, the NgR₂₇₋₃₁₁ polypeptide domain of the fusion protein would be expect to limit the myelin-associated inhibition of neurite outgrowth and promote axonal regeneration in chondroitinase digested regions of a spinal cord injury.

[0070] For cloned (NgR₂₇₋₃₁₁), the precise region of interest, or fragments can be derived from the initial clone. The biological activity of NgR₂₇₋₃₁₁, its fragments, and fusion polypeptides including NgR₂₇₋₃₁₁ may be confirmed using an in vitro assay for the collapse of growth cones on neurons. The growth cone collapse assay has been established and the addition of MAG or Nogo66 causes the collapse of growth cones in a dose dependent manner. If NgR₂₇₋₃₁₁, its fragments, and fusion polypeptides including NgR₂₇₋₃₁₁ are active is biologically active then they should inhibit the MAG and Nogo66 mediated growth cone collapse. Growth cone collapse data may be collected by the inspection of photomicrographs of DRG neurons in response to Nogo66 and MAG.

[0071] The L1 polypeptide is a member of the immunoglobulin superfamily of cell adhesion molecules and is expressed in growing axons, glial progenitor cells and Schwann cells throughout life, but has only limited expression in the CNS. L1 interacts with itself and with other extracellular molecules, such as the FGF receptor to promote neurite fasciculation and growth. Expression of L1 is generally associated with a permissive environment for axonal regeneration the, precise region of interest, or fragments of L1 can be derived from the initial clone. For example, Schwann cells that express L1 support peripheral nerve regeneration and axonal growth is observed in the optic nerves from transgenic mice that express L1 in astrocytes

but not in wild-type optic nerves. Fibroblasts engineered to express L1 support axonal growth when transplanted into spinal cord. Finally, a soluble form of L1 linked to Fc promoted functional recovery following acute SCI. If L1 is linked to proteoglycan degrading molecule like chondroitinase ABCI in a fusion protein it is reasonable to expect the fusion protein to promote axonal regeneration in chondroitinase digested regions of a spinal chord injury.

[0072] The neuregulins and their receptors comprise a diverse growth factor and receptor tyrosine kinase system that has been demonstrated to be essential for organogenesis in the CNS, muscle epithelial and other tissues. GGF2 is a soluble isoform of the neuregulin 1 gene. It was initially characterized as a Schwann cell mitogen 32, but subsequent studies have demonstrated direct actions on oligodendrocytes, neurons and other cell types. GGF2 diminishes demyelination and inflammation and enhances remyelination in a mouse model for multiple sclerosis. Based on these results, it is reasonable to expect that a recombinant human GGF2 is a potential treatment for demyelination associated with SCI. With GGF2 is linked to a proteoglycan degrading molecule like chondroitinase ABCI it is reasonable to expect to promote remyelination of axons in chondroitinase-digested regions of a SCI.

Table 2. Examples of non-limiting GGF2 fragments for expression in *E. coli* for use in compositions and fusion polypeptide compositions.

Construct	Domain	Nucleotide Seq	Amino Acid Seq	Mol. Wt.
GGF2-FL	Full length GGF2		M1-E422	46kDa
GGF2/1	Ig domain +EGF domain	748-1206	L250-C402	16.8 kDa
GGF2/2	Ig domain to C-terminus	748-1266	L250-E422	19 kDa

GGF2/3	EGF domain	1048-1206	T350-C402	5.7 kDa
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[0073] The efficacy of a composition of the present invention, such as a proteoglycan degrading molecule and a molecule that blocks the activity of an axon growth inhibitor either as a mixture or as a fusion protein can be evaluated using a validated rat SCI model at three different levels of SCI severity.

[0074] A proteoglycan degrading molecule like Chondroitinase ABCI is commercially available in small quantities as a naturally derived enzyme (Seikagaku Corporation) and can be made by a recombinant production system to have essentially the same activity as the enzyme purified from *Proteus vulgaris*. Genomic DNA can be isolated from *Proteus vulgaris* using DNeasy Tissue kit (Qiagen). PCR primers can be synthesized with an NdeI restriction site at the 5' end and a BamHI site at the 3' end having sequences 5'- CAT ATG GCC ACC AGC MT CCT GCA TTT G-3'(F2) and 5'- GGA TCC TCA AGG GAG TGG CGA GAG-3'(R) respectively, to synthesize the mature protein. The 3.0 kb PCR products can be ligated into pCR 2.1 vector (TOPO cloning kit, Invitrogen) and transformed into DH5a competent cells (Invitrogen). Plasmid DNA can be isolated from a number of clones screened by digestion with EcoRI restriction enzyme. The integrity of a gene prepared in this way can be confirmed by repeated DNA sequencing.

[0075] The chondroitinase ABCI sequence can cloned into a PET vector (Novogen) for expression in *E. Coli*. After induction of gene expression with IPTG the bacteria can lysed by sonication with the concomitant extraction of chondroitinase ABCI with Triton X-114/PBS. The inventors discovered that the majority of recombinant chondroitinase ABCI was found in the cytosolic fraction of the bacterial cell lysate that enabled the develop a chondroitinase ABCI purification protocol that yields an enzyme with high activity at high yields. The protocol

includes cation-exchange chromatography as a capture step and gel filtration as a polishing step. After these steps chondroitinase ABCI reaches a purity of ~95%. Anion exchange membrane filtration (Intercept Q, Millipore) can be used for endotoxin and host DNA removal. This step is expected to remove approximately 75% of the endotoxin. Following filtration, chondroitinase ABCI can be dialyzed into volatile buffer, pH 8.0 and lyophilized to dryness. The final product is stable at -70°C for long term storage. The purified cABCI is a highly basic protein with pI~9.5 as determined by IEF-PAGE analysis of the samples from the crude cell lysate.

[0076] A variety of analytical methods can be used to compare the enzymatic activity of the recombinant version of chondroitinase ABCI to that of a commercially available form of the enzyme (Seikagaku Corporation) purified from *Proteus vulgaris*. The methods may be adapted to evaluate the activity of fusion proteins including proteoglycan degrading polypeptides like chondroitinase. Specific activity measurements were obtained using an accepted spectrophotometric assay that measures the change in absorbance due to the production of reaction products from the degradation of proteoglycans. The recombinant form of chondroitinase ABCI had approximately 25% higher specific activity than the Seikagaku chondroitinase ABCI. Size exclusion chromatography can be used to compare the hydrodynamic properties the enzymes. The elution profiles for recombinant enzyme was identical to that of the naturally derived enzyme.

[0077] A form of zymography can used to further characterize the enzyme and may be adapted for characterization of the fusion proteins. Polyacrylamide gels can be polymerized in the presence of aggrecan, a substrate for chondroitinase ABCI. Enzyme samples may be resolved on the aggrecan-impregnated gels by electrophoresis in the presence of SDS. The gels can then be subjected to a renaturation step wherein the SDS was extracted and the enzymes

allowed to refold. Enzyme refolds and regains activity then digests aggrecan within the gel and the resulting loss of carbohydrate in that region of the gel can be visualized by a carbohydrate-specific stain. A similar loss of carbohydrate in the gel would be expected for active forms of a fusion protein including a proteoglycan degrading polypeptide portion. In the case of recombinant Chondroitinase ABCI, its activity can be visualized as a clear spot in the zymogram. The zymography results were consistent with the spectrophotometric analysis that demonstrates that the recombinant form of chondroitinase ABCI has the same or greater specific activity as the naturally occurring form.

[0078] HPLC methods may be used for detecting the four and six sulphated disaccharides (Δ 4DS and Δ 6DS, respectively) liberated as a result of chondroitinase ABCI digestion of CSPG. The two disaccharides can be effectively resolved by anion exchange chromatography. The HPLC assay has been validated by showing that the quantitation of Δ 4DS and Δ 6DS from chromatograms yields a linear relationship to the amounts injected into the HPLC. Production of Δ 4DS and Δ 6DS from CSPG digestion is directly related to the amount of chondroitinase specific activity as determined by the spectrophotometric assay described above. This assay may be used as a sensitive and accurate means to independently quantitate Δ 4DS and Δ 6DS released by chondroitinase digestion of a variety of substrates and may also be used to determine the activity of chondroitinase polypeptides in a fusion protein.

[0079] Another functional assay that can be performed to characterize proteoglycan polypeptide activity is where dorsal root ganglion (DRG) neurons are plated on aggrecan or aggrecan treated with a proteoglycan like chondroitinase ABCI. It is expected that neurons plated on aggrecan will fail to adhere to the plate and extend axons. In contrast, neurons plated on aggrecan treated with a proteoglycan degrading polypeptide like chondroitinase ABCI

in a composition or as part of a fusion polypeptide would be expected to adhere to the surface and extend axons. The extensive axon growth, which is observed for chondroitinase ABCI is believed to be due to the digestion of the carbohydrates on the aggrecan core protein which creates a more permissive substrate for axon growth.

[0080] The rat contusion model of SCI is a clinically relevant model and may be used to evaluate the efficacy of fusion proteins and other compositions of the present invention for promoting axon regeneration. With a contusion SCI, cells are destroyed, hemorrhage ensues and inflammation begins. Destroyed cells are removed by macrophages and a reactive gliosis begins. A cystic cavity is formed and the gliosis matures into a glial scar. Myelin in the area is destroyed and many local neurons that are not destroyed are left in a demyelinated state.

[0081] The forceps compression model of SCI is a contusion model developed and characterized. This model has been validated and results in injuries that are very similar to the more widely used impactor models. The model can involves a forceps compression at vertebral level T9/T10. The forceps compress the cord to a width of 0.9, 1.3 or 1.7 mm for 15 seconds. These three levels of compression allow a severe, mild or moderate injury. This model has been validated using the open field locomotor testing and the Basso, Bresnahan and Beattie (BBB) scoring system. It was also characterized histologically. Behavioral testing and BBB scoring demonstrated that the forceps produce a highly reproducible injury, with recovery similar to that seen with impactor models. These testing and scoring data demonstrate that the forceps compression model is comparable to other contusion models and sufficiently reproducible to be used as an experimental model of SCI and may be used for the evaluation of compositions of the present invention..

[0082] Tissue from forceps compression injured animals can be processed histologically to examine white matter sparing, glial scar and cyst formation. As with other contusion SCI models, a central cyst is formed after injury, with a size that increases with increased injury severity (decrease forceps gap). Around the cyst a glial scar forms that is characterized by astrogliosis (GFAP), macrophage activation and myelin wasting.

[0083] Various aspects of the present invention will be illustrated with reference to the following non-limiting examples.

EXAMPLE 1

[0084] This example illustrates how a proteoglycan degrading molecule may be administered, studied, and demonstrated to show a functional improvement in animals having a model contusion injury.

[0085] The forceps compression model of SCI is a contusion model developed and characterized. This model has been validated and results in injuries that are very similar to the more widely used impactor models. The model can involves a forceps compression at vertebral level T9/T10. The forceps compress the cord to a width of 0.9, 1.3 or 1.7 mm for 15 seconds. These three levels of compression allow a severe, mild or moderate injury.

[0086] Rats were injured with the forceps compression model at vertebral T9/T10/ At the time of surgery an intrathecal catheter was placed for delivery of chondroitinase. Animals were treated every day for one week and then on alternating days for one week with 0.06 U/ dose chondroitinase ABC I (Seikagaku), penicillinase or artificial cerebrospinal fluid (aCSF). Doses and controls were derived from Bradbury et al., 2002. Behavior was assessed using open-field locomotor testing and the BBB scoring system at day 2 and then weekly post injury for ten weeks.

[0087] FIG. 5A shown are mean BBB scores for animals treated with chondroitinase ABC I, penicillinase and aCSF. Rats treated with aCSF or penicillinase recovered to a mean BBB score of about 4. Rats treated with chondroitinase recovered to a mean BBB score of about 8. Multiple animals recovered to scores above 10, indicating supra-spinal input. The chondroitinase scores were significantly different from both control groups by ANOVA and post hoc Tukey.

[0088] Tissue from these animals was processed immunohistochemically for glial fibrillary acidic protein (GFAP) to assess general scar architecture. Tissue was also stained with a Weil stain and a silver degeneration stain to assess myelin and neuron degeneration, respectively. Interestingly no obvious differences in these parameters were noted between experimental and control treated tissues.

[0089] In FIG. 5B the large lesion comprising several segments and the sparing in the ventral cord. Weil staining revealed extensive demyelination in both the treated and untreated animals. The GFAP images (middle set) demonstrate the extent of the scar that is formed following forceps injury. The bottom amino cupric silver degeneration stain demonstrates the vast neural degeneration extending both rostrally and caudally after injury. Again, no obvious differences were noted between chondroitinase treated and control tissues.

[0090] Additional preliminary experiments have been performed at moderate injury levels and significant improvement in open-field locomotor activity was observed with chondroitinase treatment. Animals receiving chondroitinase ABCI recovered to a mean BBB score of 9.1 at ten weeks following injury, compared with 7.1 for the penicillinase controls. The consistency of the data (SEM's of 0.6 and 0.3, respectively) and the region of scores on the BBB scale make this 2 point change not only statistically significant, but also clinically meaningful. A

score of 9 indicates plantar placement with weight support or frequent to consistent weight support with dorsal stepping while a score of 7 indicates movement of each joint in hind limb but with no weight support and no consistent sweeping of limbs. An examination of the individual animal scores at ten weeks shows that 6 of 12 animals in the chondroitinase group recovered to scores of 9 or above, while only one of 12 animals in the penicillinase group recovered to a score of 9. One animal from each group was removed from analysis because its score failed to ever rise above 2.5, indicating an injury severity outside the model norms. FIG. 5C and FIG. 5D includes a scatter plot of scores at 10 weeks for each animal in the penicillinase and chondroitinase treatment groups for the moderate injury. Group means are shown below.

[0091] The results show in this well controlled study, that chondroitinase improves open-field locomotor function in Rats that were injured with the forceps compression model at vertebral T9/T10. Animals recovered to mean BBB scores of 9.7 and 9.9 for the penicillinase and chondroitinase groups, respectively. This study demonstrates a significant effect at severe and moderate injury levels, but not mild injury levels. No significant differences were noted between any groups in the mild injury study (1.3 mm forceps). It is unclear if chondroitinase is not effective with mild injury, if chondroitinase effected changes not adequately assayed by the open-field locomotor testing such as stride length, paw placement, sensory or autonomic functions. Preliminary analysis of histology from animals in this study confirmed placement of the catheters and injuries in each animal. Ongoing experiments sacrificing animals at time points after injury with and without chondroitinase treatment will characterize the CSPG content of the scar and the effects of chondroitinase digestion on stub-antigen expression. Future experiments will expand the battery of behavior tests and examine the cord for evidence of regeneration with tract tracing.

[0092] Two acute toxicity studies were conducted in rats. The first was an intravenous (IV) study wherein rats were injected with 0, 0.2, 0.775 or 7.775 mg/kg chondroitinase ABCI. In the second study, intrathecal (IT) catheters were placed over the spinal cords using the same methods employed in the SCI animal studies and 0.06, 0.6 and 6.0 units of chondroitinase ABCI was delivered through the IT catheters. These doses were 1, 10, and 100-fold greater than the estimated local concentrations of chondroitinase ABCI achieved with IT catheters in the SCI experiments. Animals were monitored for pain and distress and body weights were acquired daily. No overt reactions were observed during or immediately after chondroitinase ABCI dosing. No swelling, inflammation, bruising or necrosis was noted at the injection site for the IV experiment. No changes in body temperature were observed in animals treated via the IT route. No alterations in feeding, grooming or vocalizations were noted. Animals were assessed for motor behavior in an open pool. No abnormalities were noted by the animal care staff or behavioral specialists. Animals displayed no signs of joint tenderness or swelling. There were no significant differences in weight change between the treatment groups. The results demonstrate that chondroitinase ABCI treatment is not associated with acute toxicity using IV and IT doses substantially greater than the efficacious IT doses.

EXAMPLE 2

[0093] This example describes the preparation of Nogo-receptor agonist, L1 neural cell adhesion protein, and GGF2 polypeptide domains which can be used for compositions and fusion proteins of the present invention.

[0094] A soluble portion of the human Nogo receptor spanning amino acids 27 to 311 (NgR₂₇₋₃₁₁) was selected as it has been shown to inhibit the binding of Nogo66, MAG, and MOG to membrane-bound NgR. Primers were designed flanking this region, and RT-PCR was

performed using human hippocampal RNA (BD Biosciences). The 1.05kb region was successfully amplified and purified.

[0095] L1 was prepared through a CHO cell line from the laboratory of Dr. Melitta Schachner that secretes human L1 as a fusion protein with human Fc (L1-Fc). The cells were grown in roller bottles and then L1-Fc was purified from the conditioned media using protein A affinity column chromatography. The purity of L1-Fc was assessed by SDS-PAGE and a single band at the appropriate molecular weight was observed. The biological activity of L1-Fc was confirmed using a neurite outgrowth assay. Tissue culture plates were coated with either poly L-Lysine or L1-Fc and then cerebellar granule cells from postnatal day 10 rats were isolated and placed into culture on the substrates. It was observed that neurons plated on the L1-Fc substrate exhibited a substantial number of long neuritis compared to the polylysine substrate controls. These results demonstrate that the L1-Fc produced is biologically active in promoting neurite outgrowth. This neurite outgrowth assay will be used to assess the biological activity related to the L1 portion of the chondroitinase ABCI fusion protein..

[0096] A CHO cell line that secretes a soluble form of full length GGF2 glycoprotein was obtained from CeNes. Extensive optimization of media and purification methods was completed to obtain essentially pure and biologically active GGF2. A number of analytical methods were developed to characterize the GGF2 including SDS-PAGE, isoelectric focusing, peptide mapping and carbohydrate analysis. For example an SDS-PAGE gel of reduced and non-reduced GGF2; the isolate is essentially free of contaminating proteins and shows the expected molecular weight and monomeric structure. The biological activity of GGF2 was assayed using a primary rat Schwann cell proliferation method and the expected effect was reproducibly obtained with four independent batches of GGF2. Another functional assay was

developed that measures the phosphorylation of Akt kinase, a downstream cell signaling component of the erbB receptor pathway. It was observed that there is a dose dependent phosphorylation of Akt kinase

EXAMPLE 3

[0097] This example illustrates the manipulation of chondroitinase ABCI for construction of chimeras.

[0098] Chondroitinase ABCI was cloned from *P.vulgans* and expressed in *E.coli*. Surprisingly repeated attempts to create N-terminal fusion proteins were not successful because the N-Terminal fusion portion was cleaved from chondroitinase ABCI during synthesis. However, catalytically active deletion mutants with N-terminal His fusion tags designated ABC I-NΔ2O, ABC I-NΔ4O and ABCI-NΔ6O were prepared by deleting 20, 40 and 60 amino acids respectively from the N-terminus of the mature ABCI protein. Unlike the full length chondroitinase ABCI, the N-terminal deletion mutants are capable of synthesizing a 6xHis tag as an N-terminal fusion protein. It was also observed that deletion of 80 amino acids from the C-terminal end formed a mutant with proteoglycan degrading activity of chondroitinase ABCI as tested in a zymography assay.

[0099] Fusion proteins with NgR₂₇₋₃₁₁ or L1 with a proteoglycan degrading molecule such as chondroitinase ABCI will utilize mammalian expression and can be performed in CHO cells. The cDNA for chondroitinase ABCI has been cloned in pSECTag vector (Invitrogen) in the proper reading frame. A CHO cell line having secretary chondroitinase ABCI can be developed and the conditioned medium can be tested for catalytic activity by zymography assay to confirm that chondroitinase ABCI expressed in mammalian cells is functional. A mammalian

cell codon optimized version of chondroitinase ABCI can be synthesized by methods known in the art for use for CHO cell expression.

[00100] GGF2 is a spliced variant of the NRG1 gene expressed in brain and spinal cord of adult humans. It is a glycosylated protein of molecular mass between 66-90 kDa. The inventors have discovered that recombinant GGF2 expressed in CHO cells is highly glycosylated and promotes Schwann cell proliferation in vitro and further that an EGF-like domain of NRG1 expressed in *E.coli* is fully functional in promoting myocyte proliferation and survival.

[00101] The inventors have expressed fragments of GGF2 in *E.coli* as described in the preliminary data section. The specific GGF2 domain responsible for Schwann cell proliferation and thereby remyelination can be determined. If the Ig and EGF domains together or separately show biological activity in vitro, then they can be used to form chimeric fusion proteins.

[00102] Cloning and expression of NgR₂₇₋₃₁₁ in CHO cells. An NgR fragment corresponding to residues 1-359 was isolated by RT-PCR from human hippocampus poly K RNA (BD Biosciences) and its structure can be confirmed by DNA sequencing. The gene fragment corresponding to residues 27-311 can be cloned from the larger fragment and then subcloned in pSECTag vector (Invitrogen) in the proper reading frame to express the fragment as a secretary protein in CHO cells. The plasmid DNA containing the NgR₂₇₋₃₁₁ gene can be transfected in CHO cells and the cell line producing NgR₂₇₋₃₁₁ can be selected under the selection pressure of hygromycin B. An NgR₂₇₋₃₁₁-ABCI chimera expression plasmid can be constructed for expression in a CHO cell expression system using methods known in the art.

EXAMPLE 4

[00103] This example illustrates methods which may be used to purify and isolate expressed chondroitinase ABCI, and GGF2 domains expressed in *E. coli*. These method may be applied to purification of chimeric fusion proteins of the present invention.

[00104] An efficient *E.coli* recombinant expression system and a purification process for chondroitinase ABCI have been developed by the inventors that could be applied for the purification of chondroitinase ABCI chimeric derivatives.

[00105] For example, the expression of various GGF2 domains in the chondroitinase ABCI expression host, *E.Coli*, has been preformed and the following peptides have been tested: aa250-402, aa250-422 and aa350-402. These expressed peptides were found in the soluble fraction of the *E.coli* lysates. It is reasonable to expect that final chimeric products of these peptides with chondroitinase ABCI in *E.coli* will also be soluble. In addition, it is expected that the charge characteristics of the chimeric products when compared to the recombinant chondroitinase ABCI will be similar. Thus, the theoretical isoelectric point (p1) values for the GGF2 peptides are 9.3 for aa250-402, 9.18 for aa250-422 and 7.55 for aa350-402. Fusing the first two peptides with chondroitinase ABCI is expected to result in chimeric proteins with p1 values still above 9. In this case SP chromatography is expected to perform well as the capturing step. The smallest GGF2 peptide, aa 350-402, will reduce the p1 of the final chondroitinase ABCI chimera to approximately 8.4. This change may require optimization of the capture step conditions.

[00106] The chimeric products of chondroitinase ABCI with NgR₂₇₋₃₁₁ and L1 peptides can be expressed in a CHO cell system. Prior to purification of the expressed chimeric proteins, the growth conditions for the cell line producing either the NgR₂₇₋₃₁₁ or L1 chimeric proteins will be optimized in serum-free medium. A detailed media optimization study can be

performed to determine the highest production conditions. Scale-up volume can be decided based on the rate of production of the chimeric proteins (pg/cell/day). Conditioned media from the various chimera-producing cell lines can be collected and subjected to tangential flow filtration. Ion-exchange chromatography can be used for capturing the secreted proteins from conditioned media, gel filtration chromatography can be used as a polishing purification step and then anion-exchange membrane filtration can be used for endotoxin and DNA removal. At each step the efficiency of the purification will be analyzed by SDS-PAGE, and spectrophotometric quantitation of protein concentration.

EXAMPLE 5

[00107] This prophetic example describes in vitro assessment of chimera biological activity: Each chimera can be assayed for chondroitinase enzymatic activity and the specific biological activity of each fusion partner.

[00108] The first step in analysis may employ conventional protein biochemical methodologies to confirm the fidelity of gene expression. These include SDSPAGE, IEE, mass spectrometry and size exclusion chromatography.

[00109] Chondroitinase chimera specific activity can be determined using a standard and uniformly accepted spectrophotometric assay. The production of reaction products from the catalytic activity of a chondroitinase chimeric polypeptide can be determined by a measurement of the absorbance of the product at a wavelength of 232 nm. A typical reaction mixture consists of 120 microliters of reaction mixture (40mM Tris, pH 8.0, 40mM NaAcetate, 0.002% casein) combined with substrate (5 microliters of 50 mM chondroitin C) and 1.5 microliters of chondroitinase ABCI chimeric fusion polypeptide or test sample. The change in absorbance units is the initial rate which can be converted to a unit activity measurement.

[00110] Disaccharide HPLC assay: The catalytic activity of chondroitinase

chimeric polypeptide on a CSPG substrate is expected to releases two species of sulphated disaccharides including α - Δ AUA-[1→3]-GalNAc-4S (Δ 4DS) and α - Δ AUA-[1→3]-GalNAc-6S (Δ 6DS). These species can be resolved by HPLC and the quantitation from the resulting chromatograms is a sensitive and accurate measure of chondroitinase activity. To perform the assay, samples from chondroitinase digestion reactions carried out with wild-type chondroitinase and chondroitinase fusion proteins can be clarified by centrifugation and then subjected to a validated anion exchange HPLC method as follows. A Dionex CarboPac PA-10 analytical column (4 x 250mm) fitted with a Dionex CarboPac PA-10 (4x 50mm) guard column can be used with a mobile phase consisting of a gradient of water at pH 3.5 (Buffer A) and 2M NaCl, at pH 3.5 (Buffer B). Detection may be set at a wavelength of 232 nm. A flow rate of 1 mL/minute and a 45 minute continuous gradient of 100% A to 100% B affords acceptable resolution of Δ 4DS and Δ 6DS. Standard curves can be generated using known amounts of Δ 4DS and Δ 6DS.

[00111] Zymography allows resolution of proteins by molecular weight with a

concomitant assessment of chondroitinase activity. A 10% polyacrylamide gel may be polymerized in the presence of 85pg/ml of aggrecan. Samples can be boiled in an SDS-loading buffer and then the analytes can be resolved by electrophoresis. After separation the gel can be incubated for 1 hour at room temperature in 2.5% Triton X100 then 16 hours at 37°C in fresh 2.5% Triton X-100. During these incubations, the SOS can be extracted from the gel and the chondroitinase refolds and digests the aggrecan in its immediate vicinity. After the refolding process the gel can be stained for carbohydrate: The gel can be first incubated in 0.2% Cetylpyridinium for 90 minutes at room temperature and then transferred into 0.2% Toluidine

Blue in 49:50:1 H₂O, Ethanol, and Acetic Acid for 30 minutes. The gel can then be fully destained. Following destaining the gel can be incubated overnight in a 50 microgram/ml solution of Stains-All (Sigma) in 50% ethanol. Chondroitinase activity can be detected as a clear spot in the gel that is coincident with the molecular weight of the enzyme. The size of the clearing has been shown to be almost linearly related to Unit activity.

[00112] The NgR₂₇₋₃₁₁-chondroitinase chimera can be assessed for the activity of the decoy Nogo receptor. The assay can be used to measures the collapse of growth cones: Dorsal root ganglia (DRG) are dissected from postnatal day 1 (P1) Sprague Dawley rat pups and dissociated in 200 U/ml collagenase I (Worthington) and 2.5 U/ml dispase (Boehringer/Roche) 2 times for 30 min. at 37°C. Enzymes are removed and DNase (0.5mg/ml) can be added to the ganglia. Trituration may be done with a pipette tip attached to a 1000 µl Pipetman. The resulting cell suspension can be filtered through a 40 micron cell filter and centrifuged at 70xg for 5 min. Cells can be re-suspended in DMEM/10% FBS and pre-plated for 2 hours on a non-coated tissue culture plate (100mm diameter). Non-adherent neurons are removed and plated at 10,000 cells/well in a Poly-lysine/laminin-coated 24 well plate in serum-free Neurobasal/B27 with 50ng/ml NGF. After 20 to 24 hours, MAG or Nogo66 can be added at varying concentrations for 1 hour at 37°C to induce growth cone collapse. NgR₂₇₋₃₁₁-chondroitinase chimera can be added at various concentrations to compete with MAG and Nogo66 and thus protect the neurons from growth cone collapse. Cultures may be fixed by adding an equal volume of pre-warmed 8% paraformaldehyde/0.6 M sucrose to the medium for 20 min. while the cells are kept on a 37°C hot plate. Growth cones may be labeled with AlexaS68 phalloidin (Molecular Probes). Briefly, cells can be permeabilized with 0.1% Triton-X 100 for 5 min. at RT, blocked for 20min. in 1% BSA in PBS and incubated in phalloidin, diluted 1:40 in 1% BSA,

in PBS for 20mm at RT. Cells can be washed in PBS and mounted in Fluorescent Mounting Medium (DAKO). The percentage of collapsed growth cones is determined by analyzing a minimum of 100 growth cones per well under a 40x objective.

[00113] The biological activity L1 fusion proteins may be determined using a standardized neurite outgrowth assay. After etching a 25mm circle in the center of a tissue culture-treated 35mm dish, 1 ml of 10 µg/ml poly-lysine can be added to each etched circle and incubated at 37°C for 60 minutes. The poly-lysine provides a negative control for neurite outgrowth. Etched circles may be washed 2 times with Hank's balanced salt solution plus calcium and magnesium (HBSS⁺⁺) after the last rinse; 1.2 µl of L1-Fc (0.6µM, 0.3µM, 0.15 µM, 0.075 µM) is spotted onto the petri dishes to serve as a positive control and various concentrations of the L1-chondroitinase fusion protein can be applied to the plates as test samples. The plates can be incubated for 1 hour at room temperature. The spots are lightly aspirated, so as not to dry them out, and immediately washed 2 times with 1 ml HBSS⁺⁺ and then 1 ml of 1% BSA in PBS is added to each etched circle. After about 15 minutes at room temperature the etched circles can be washed 2 times with HBSS⁺⁺, and once with bioassay medium (NeuralBasal(Gibco) + B27 supplements (Gibco) + L-glutamine + L- glutamic acid, penicillin and streptomycin (100 U/ml) + 1% fetal bovine serum which will remain on the petri dishes until the time of the assay. To provide neurons to plate on the substrates Cerebellar granule cells from postnatal day (PND) 9 or 10 are harvested. The brain is removed from the cranium of 1 pup, and the cerebellum separated from the rest of the tissue. Meninges is removed, and the cerebellum is placed in ice cold HBSS⁺⁺. The tissue is minced and trypsinized using 0.25% trypsin for 15 minutes at 37°C. Trypsin action is inhibited by adding 0.5mg/ml soybean trypsin inhibitor (Gibco). The tissue is rinsed with HBSS⁺⁺ and triturated using a flame

narrowed Pasteur pipette coated with EBS. Dissociated cells are pelleted at 500xg for 3 minutes, the supernatant is decanted and the cells are resuspended in 2ml of growth medium. After lightly triturating, the cell suspension is carefully layered on top of a 3.5%BSA (in PBS) cushion, and spun at 500xg for 3 minutes. Supernatant is aspirated and the pellet is resuspended in 2 ml growth media. A cell count is performed and the cells are diluted to a final working concentration of 1.5×10^5 cell/ml. A 300 μ l aliquot of diluted cells is added to the center of each plate, resulting in the even distribution of 6×10^4 cells across the entire surface area of the etched circle. The cells are allowed to grow for 16 hours at 37°C in a humidified environment supplemented with 5% CO₂. The next day, plates are removed from the 37°C incubator, cells are fixed with 4% paraformaldehyde and the outgrowth of neurites is recorded by photomicroscopy.

[00114] Biological activity assays for GGF2 may be performed using Schwann cell proliferation. Sciatic nerves are dissected from three day old Sprague Dawley rat pups and dissociated in L-15 medium (Invitrogen) containing 0.25% trypsin and 0.03% collagenase type I (SIGMA) for 15 minutes at 37°C. Nerves are centrifuged at 400xg for 5 minutes and dissociation medium is replaced by DMEM/10% FBS. Nerves are triturated using a 10 ml syringe with a 21g needle and subsequently a 23g needle. The cell suspension is filtered through a 40 μ m mesh placed over the opening of a 50ml conical tube. Cells can be centrifuged at 400xg for 5min, plated in a poly-D-Lysine (PDL)coated T-75 flask at a density of approximately 5 million cells in 15 ml of DMEM/10% FBS/PenStrep and incubated in a 37°C incubator regulated with 10% CO₂. After a 24 hour incubation the cells can be washed twice with DMEM/10% FBS and re-fed with fibroblast inhibition medium consisting of DMEM/10%FBS and 10 μ l/ml of 1 mM cytosine arabinoside (Ara-C). After an incubation time of 2 to 3 days the fibroblast inhibition medium is replaced with Schwann cell growth medium (DMEM/10% FBS/ 150 ng/ml

of GGF2/5 μ M forskolin). Cells can be expanded and aliquots of 2×10^6 cells/ml are frozen in DMEM/10%DMSO, 54% FBS in liquid nitrogen. At the time of use, Schwann cells are thawed and plated at a density of about 16,000 cells/well in a PDL-coated 96 well tissue culture plate in DMEM/5% FBS. After about 24 hours GGF2 is added in serial dilution ranging from 100 ng/ml to 0.78ng/ml containing 5 μ M forskolin to establish a standard curve. Samples of GGF2 fusion proteins can also be added in serial dilution together with 5 μ M forskolin. BrdU is added at 10 μ M. Cells are incubated for 48 hours in a 10% CO₂ incubator. A BrdU ELISA kit from Roche Applied Science can be used (Cat. No. 1 647 229) to detect Schwann cell proliferation. Medium is poured out of the plate and plate is tapped on tissue paper. A 200 μ l/well aliquot of Fix/Denat is added and incubated for 30 min at 15-25°C. A 100 μ l/well aliquot of anti-BrdU-POD working solution (lyophilized antibody is dissolved in 1.1 ml double distilled water and diluted 1:100 with antibody dilution solution) can be added and incubated for 90 min at room temperature. Wells may be washed 3 times with 200-300 μ l/well of washing solution. A 100 μ l aliquot of substrate solution is added and incubated for 15 minutes or until color development is sufficient for photometric detection. The Plates are read on a SpectraMax plate reader at 450nm either before addition of stop solution at 370nm or after addition of 50 m1 1N sulfuric acid to each well.

[00115] Akt [pS473] ELISA: C6 glioma cells, obtained from ATCC, are grown in DMEM/10%FBS in a T-75 flask to confluence. After trypsinization, cells are plated in a 24 well plate at a density of 500,000 cells/well in 0.5 m1 of medium. One day after plating the cells are treated with GGF2 (batch: Glu from Lonza Biologics) at serial dilutions ranging from 0.78 to 100 ng/ml for 30 minutes at 37°C to establish a standard curve. As a negative control, wortmannin, a PI3-kinase inhibitor, is added to cells at 10 nM for 30 minutes before the

addition of GGF2. Samples of GGF2 fusion proteins will also be added in serial dilution. The cells are washed with PBS and then extracted with 100 μ l of cell extraction buffer (10 mM Tris, pH 7.4, 100 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1 mM NaF, 20 mM Na₄P₂O₇, 2 mM Na₃VO₄, 1% Triton X-100, 10% glycerol, 0.1% SDS, 0.5% deoxycholate, 1 mM PMSF, protease inhibitor cocktail (Pierce)). Cell extracts are kept at -80°C until further use. An ELISA kit from Biosource (Cat. #KHO01 II) can be used to measure phosphorylated Akt kinase levels. Briefly, 100 μ l of a 1:50 dilution of samples and a serial dilution of Akt [pS473] standards are loaded into wells pre-coated with anti-Akt antibody for 2 h at room temperature (RT) or overnight at 4°C. Wells are washed and anti-Akt [pS473] antibody is added and incubated for 1 hour. After washing, HRP-conjugated anti-rabbit antiserum is added to wells and incubated for 30 min. After washing, TMB chromogen is added to wells and incubated for 30 min at RT before the reaction is stopped with stop solution. The plate is read on a SpectraMax plate reader at 450 nm. The optical densities are plotted against the Akt [pS473] standards and the concentration of pAkt in the C6 samples are deduced from the standard curve.

EXAMPLE 6

[00116] This example illustrates construction of a fusion protein of chondroitinase polypeptide and a TAT cellular transduction peptide.

[00117] The gene sequence encoding a chondroitinase enzyme can be functionally linked to the protein transduction domain from HIV called the TAT Peptide. The resultant chimeric genes TAT-chondroitinase ABCI fusion DNA construct are shown in FIG. 1. It was observed that during bacterial expression of this construct, the TAT peptide was removed from

the chondroitinase enzyme at some processing point during the bacterial growth. The removal of n-terminal linked peptides was also observed during expression of an n-terminal histidine-tagged chondroitinase ABCI enzyme.

[00118] Deletion mutants of the chondroitinase ABCI enzyme were generated where the mutant was missing a certain number of amino acids from the n-terminal portion of the polypeptide but maintained its proteoglycan degrading activity. It was observed that these n-terminal deletion maintained a histidine-tag that was attached to the n-terminus. It is expected that various TAT-deletion mutant chondroitinase ABCI fusion DNA construct can be expressed without removal of the TAT polypeptide during expression. For example, the TAT peptide may be fused at the N-terminus of deletion mutants like ABCI-NΔ20 or ABCI-NΔ60 deletion mutant. Without wishing to be bound by theory, the inventors believe that the native proteoglycan degrading enzyme contains a signal sequence that is attached to the n-terminus. This signal sequence is removed during the natural production in bacteria and in production of the cloned enzyme in *E. coli*. It is thought that some signal within the n-terminal amino acids instructs the bacteria to remove anything attached to this end.

[00119] A DNA construct with the TAT-peptide attached to the N-terminus of one of the chondroitinase deletion mutants can be made and Western blot and protein gel showing this expressed protein and activity.

EXAMPLE 7

[00120] This example illustrates the diffusion of molecules into cells and tissue using a proteoglycan degrading composition.

[00121] A brain from an adult Sprague Dawley rat was removed from the skull and quartered into Right frontal, left frontal, right rear and left rear sections, corresponding roughly

to the frontal (front) and occipito-parietal (rear) lobes. **(A)** Right frontal quarter was placed in artificial cerebrospinal fluid (Catalog # 59-7316; Harvard Apparatus, Holliston, MA) containing the beta-galactosidase enzyme (Catalog #, G 5160; Sigma, St. Louis, MO) and chondroitinase ABC I at 0.5 U/ml (Catalog# C 3667; Sigma, St. Louis, MO) for 2 hours at 37°C. Brain quarter was rinsed several times in phosphate buffered saline (PBS) and then processed with a Beta-Gal staining kit (Catalog # Gal-S; Sigma, St. Louis, MO). The substrate-enzyme reaction (blue product) was allowed to develop for 1 hour, and the brain was rinsed several times in PBS and slabs from the middle of each brain block cut using parallel straight razors. **(B)** Left frontal quarter was placed in artificial cerebrospinal fluid (Catalog # 59-7316; Harvard Apparatus, Holliston, MA) containing the beta-galactosidase enzyme (Catalog #, G 5160; Sigma, St. Louis, MO) for 2 hours at 37°C. Brain quarter was rinsed several times in phosphate buffered saline (PBS) and then processed with a Beta-Gal staining kit (Catalog # Gal-S; Sigma, St. Louis, MO). The substrate-enzyme reaction (blue product) was allowed to develop for 1 hour, and the brain was rinsed several times in PBS and slabs from the middle of each brain block cut using parallel straight razors. **(C)** Right frontal quarter was placed in artificial cerebrospinal fluid (Catalog # 59-7316; Harvard Apparatus, Holliston, MA) containing the beta-galactosidase enzyme (Catalog #, G 5160; Sigma, St. Louis, MO) and chondroitinase ABC I at 0.5 U/ml (Catalog# C 3667; Sigma, St. Louis, MO) for 2 hours at 37°C. Brain quarter was rinsed several times in phosphate buffered saline (PBS) and then processed with a Beta-Gal staining kit (Catalog # Gal-S; Sigma, St. Louis, MO). The substrate-enzyme reaction (blue product) was allowed to develop for 1 hour, and the brain was rinsed several times in PBS and slabs from the middle of each brain block cut using parallel straight razors. **(D)** Left rear quarter was placed in artificial cerebrospinal fluid (Catalog # 59-7316; Harvard Apparatus, Holliston, MA) containing the beta-

galactosidase enzyme (Catalog #, G 5160; Sigma, St. Louis, MO) for 2 hours at 37°C. Brain quarter was rinsed several times in phosphate buffered saline (PBS) and then processed with a Beta-Gal staining kit (Catalog # Gal-S; Sigma, St. Louis, MO). The substrate-enzyme reaction (blue product) was allowed to develop for 1 hour, and the brain was rinsed several times in PBS and slabs from the middle of each brain block cut using parallel straight razors. Images of brains were acquired on a scanner and are shown in FIG. 2(I)(A-D).

[00122] Adult rat brain hemispheres were soaked in buffer alone or containing 33U/ml Chondroitinase ABCI (Acorda) for 2 hours at 37 degrees C. Hemispheres were rinsed and immediately placed in Eosin Y (Sigma) or a saturated solution of Congo Red (Sigma) in 70% ethanol. Slabs of tissue were cut and images were acquired on a scanner. See FIG. 2(II) Eosin and FIG. 2(III) Congo red.

[00123] FIG. 2(III) saturated solution of Congo Red demonstrates greater penetration through the cortex of a Chondroitinase treated brain hemisphere as compared to untreated brain. Congo Red is a negatively charged dye of 697 kDa. FIG. 2(II) Eosin Y penetration through the cortex of a Chondroitinase treated brain hemisphere looks slightly more diffuse, but penetration does not seem to be any deeper as compared to untreated brain. Eosin Y is zwitterionic, having an overall negative charged at the low pH it was used at, and it is 692 kDa. FIG. 2(I) Lobes from adult rat were incubated in beta-galactosidase alone (B&D), or with the addition of Chondroitinase ABCI (A, 0.5U/ml or C, 0.005U/m).

[00124] The results showed that chondroitinase treated tissue affected penetration of beta-galactosidase into CNS tissue. Chondroitinase had dramatic effects on the rate of Congo Red dye penetration but apparently not Eosin.

EXAMPLE 8

[00125] This example illustrates a Chondroitinase ABCI Assay Protocol which may be modified to measure the activity of Chondroitinase ABCI deletion mutant-fusion proteins or the activity of other proteoglycan degrading polypeptide-fusion proteins of the present invention.

[00126] The production of reaction products from the catalytic activity of a proteoglycan degrading molecule or fusion protein is determined by a measurement of the absorbance of the product at a wavelength of 232 nm. A typical reaction mixture consists of 120 µl of reaction mixture (40mM Tris, pH 8.0, 40mM NaAcetate, 0.002% casein) combined with a substrate (5 µl of 50 mM chondroitin C) and 1.5 µl of chondroitinase ABCI or a fusion protein of a deletion mutant of chondroitinase ABCI-TAT. Seikagaku's cABCI reference: frozen at -20°C in 5ml aliquots, use 1ml per reaction, 50 mM chondroitin C (MW 521) in water: frozen at -20°C in 100ml aliquots. Reaction mixture aliquots of about 120 µl can be prepared at 37°C for 3 min or longer. A wavelength of 232 nm, is used with the spectrometer.

[00127] Knowing the extinction coefficient for the reaction product, measuring the change in the absorbance of the reaction product at 232 nm reading over time upon addition of a known amount of the Chondroitinase ABCI or other other proteoglycan degrading polypeptide-fusion proteins to the 120 µl reaction mixture with 0.002% casein and chondroitin substrate added, the specific activity in µmol/min/mg of the proteoglycan degrading fusion protein can be determined. Seikagaku Chondroitinase ABCI has a specific activity under these assay conditions of about 450 µmole/min/mg.

[00128] Although the present invention has been described in considerable detail with reference to certain preferred embodiments thereof, other versions are possible. Therefore

the spirit and scope of the appended claims should not be limited to the description and the preferred versions contain within this specification.

CLAIMS

What is claimed is:

1. A composition comprising:
 - an isolated nucleic acid comprising a sequence that encodes a polypeptide, said polypeptide comprising at least two of: a protein transduction domain, a proteoglycan degrading domain, or a domain that promotes neural regeneration.
2. The composition of claim 1 wherein nucleic acid encode for the protein transduction domain TAT domain.
3. The composition of claim 1 wherein the any two domains of said polypeptide are bonded together by a linker polypeptide domain.
4. The composition of claim 1 wherein the nucleic acid encodes for a proteoglycan degrading domain that is a deletion mutant of chondroitinase ABCI.
5. The composition of claim 1 wherein the domain that promotes neural regeneration is L1, a variant that is at least 80% L1, or a deletion mutant of L1.
6. The composition of claim 1 wherein the domain that promotes neural regeneration is GGF2, a functional variant that is at least 80% GGF2, or a functional deletion mutant of GGF2.
7. The composition of claim 1 wherein the domain that promotes neural regeneration is NgR₂₇₋₃₁₁, a functional variant that is at least 80% homologus with NgR₂₇₋₃₁₁, or a functional deletion mutant of NgR₂₇₋₃₁₁.
8. The composition of claim 1 expressed from genetically modified cells.
9. The composition of claim 1 and cells from the CNS.
10. A composition comprising:

a polypeptide comprising at least two of: a protein transduction domain, a proteoglycan degrading domain, or a domain that promotes neural regeneration.

11. The composition of claim 10 wherein the protein transduction domain is a TAT domain.
12. The composition of claim 10 wherein two domains of said polypeptide are bonded together by a linker polypeptide domain.
13. The composition of claim 10 wherein the proteoglycan degrading domain is a deletion mutant of chondroitinase ABCI.
14. The composition of claim 10 wherein the domain that promotes neural regeneration is L1, a variant that is at least 80% L1, or a deletion mutant of L1.
15. The composition of claim 10 wherein the domain that promotes neural regeneration is GGF2, a functional variant that is at least 80% GGF2, or a functional deletion mutant of GGF2.
16. The composition of claim 10 wherein the domain that promotes neural regeneration is NgR₂₇₋₃₁₁, a functional variant that is at least 80% homologous with NgR₂₇₋₃₁₁, or a functional deletion mutant of NgR₂₇₋₃₁₁.
17. The composition of claim 10 having a chondroitinase.
18. The composition of claim 10 and a pharmaceutically acceptable excipient.
19. The composition of claim 10 and cells from the CNS.
20. A method of promoting recovery of neurological function comprising:
 - identifying a patient having comprised neurological function characterized by an SCI injury; and
 - administering a composition comprising at least two of: a protein transduction polypeptide, a proteoglycan degrading polypeptide, or a polypeptide that promotes neural regeneration, to a patient in need of such treatment.

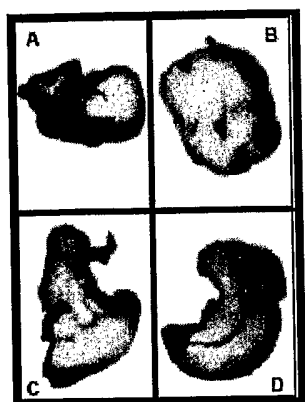
21. The method of claim 20 wherein the composition is secreted by genetically modified cells that are implanted near the site of the CNS injury.
22. The composition of claim 20 wherein the proteoglycan degrading protein is Chondroitinase ABC I, Chondroitinase ABC II, Chondroitinase AC, Chondroitinase B, Hyaluronidase 1, Hyaluronidase 2, Hyaluronidase 3, Hyaluronidase 4, PH20, or combination of these.
23. The composition of claim 20 wherein the polypeptides are linked together.
24. A method of facilitating diffusion of an agent in a central nervous system comprising:
 - administering the agent to the central nervous system; and
 - modifying an extracellular matrix in the central nervous system to allow penetration of the agent into the extracellular matrix.
25. The method according to claim 24, further comprising the step of utilizing at least one enzyme capable of cleaving proteoglycans in the extracellular matrix.
26. The method according to claim 24, wherein the proteoglycans are chondroitin sulfate proteoglycans.
27. The method according to claim 24, wherein the at least one enzyme is selected from the group consisting of chondroitinase ABC Type I, chondroitinase ABC Type II, chondroitinase AC, chondroitinase B, hyaluronidase 1, hyaluronidase 2, hyaluronidase 3, hyaluronidase 4, cathepsins, ADAMT's, PH20, and any combination thereof.
28. The method according to claim 24, wherein the at least one enzyme is administered with one or more pharmaceutical carriers.
29. The method according to claim 24, wherein the agent is selected from the group consisting of complex biologics, small molecules, and transplanted cells.

30. The method according to claim 24, wherein administering the agent is accomplished topically, locally or systemically.
31. The method according to claim 24, wherein administering the agent is by bolus injection, intravenous delivery, continuous infusion, sustained release from implants, or sustained release pharmaceuticals.
32. A method of penetration of an agent through an extracellular matrix, comprising:
administering at least one enzyme capable of cleaving chondroitin sulfate proteoglycans.
33. The method according to claim 32, wherein the at least one enzyme is selected from the group consisting of chondroitinase ABC Type I, chondroitinase ABC Type II, chondroitinase AC, chondroitinase B, hyaluronidase 1, hyaluronidase 2, hyaluronidase 3, hyaluronidase 4, cathepsins, ADAMT's, PH20, and combination thereof.
34. The method according to claim 32, wherein the at least one enzyme is administered in an effective amount.
35. The method according to claim 32, wherein the agent is selected from the group consisting of complex biologics, small molecules, and transplanted cells.
36. The method according to claim 32, further comprising the step of administering the agent topically, locally or systemically.
37. The method according to claim 32, further comprising the step of administering the agent by bolus injection, intravenous delivery, continuous infusion, sustained release from implants, or sustained release pharmaceuticals.
38. A method of enhancing uptake of an agent in a central nervous system, comprising:
administering the agent and at least one enzyme capable of cleaving chondroitin sulfate proteoglycans.

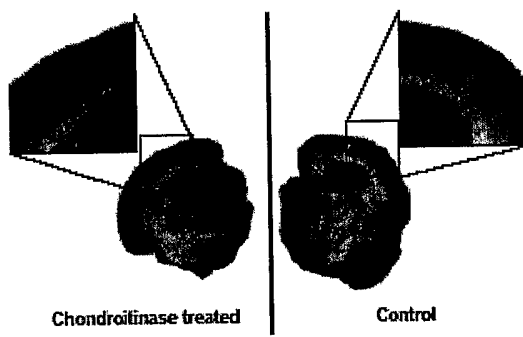
39. The method according to claim 38, wherein the at least one enzyme is selected from the group consisting of chondroitinase ABC Type I, chondroitinase ABC Type II, chondroitinase AC, chondroitinase B, hyaluronidase 1, hyaluronidase 2, hyaluronidase 3, hyaluronidase 4, cathepsins, ADAMT's, PH20, and any combination thereof.
40. The method according to claim 38, wherein the at least one enzyme is administered in an effective amount.
41. The method according to claim 38, wherein the at least one enzyme is administered with one or more pharmaceutical carriers.
42. The method of claim 38, wherein the pharmaceutical carrier is selected from the group consisting of diluents, binders, lubricants, coloring agents, disintegrating agents, buffer agents, isotonizing agents, polyols, preservatives, and anesthetics.
43. The method according to claim 38, wherein the agent is selected from the group consisting of complex biologics, small molecules, and transplanted cells.
44. The method according to claim 38, wherein administering the agent is accomplished topically, locally or systemically.
45. The method according to claim 38, wherein administering the agent is by bolus injection, intravenous delivery, continuous infusion, sustained release from implants, or sustained release pharmaceuticals.

TAT fusion chondroitinase ABCI nucleic acid

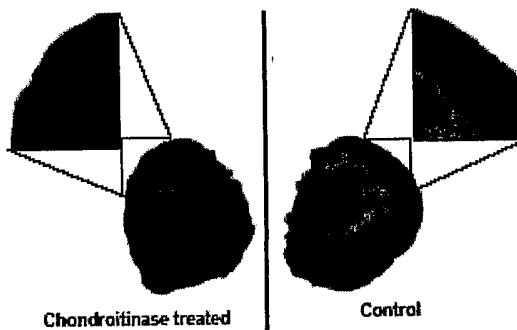
FIG. 1



(I)

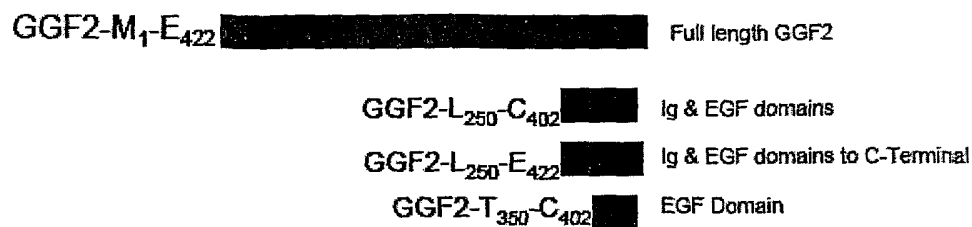


(II)



(III)

FIG. 2



(A)

N-terminal Fusion Chimera between Chondroitinase ABCI and GGF2

N	GGF2-FL	C	N	ABCI-NΔ60-CΔ80	C
	GGF2-L250-C402		N	ABCI-NΔ60-CΔ80	C
	GGF2-L250-E422		N	ABCI-NΔ60-CΔ80	C
	GGF2-T350-C402		N	ABCI-NΔ60-CΔ80	C

C-terminal Fusion Chimera between Chondroitinase ABCI and GGF2

ABCI-FL	C	N	GGF2-FL	C
N	ABCI-FL	C	GGF2-L250-C402	
N	ABCI-FL	C	GGF2-L250-E422	
N	ABCI-FL	C	GGF2-T350-C402	

(B)

FIG. 3

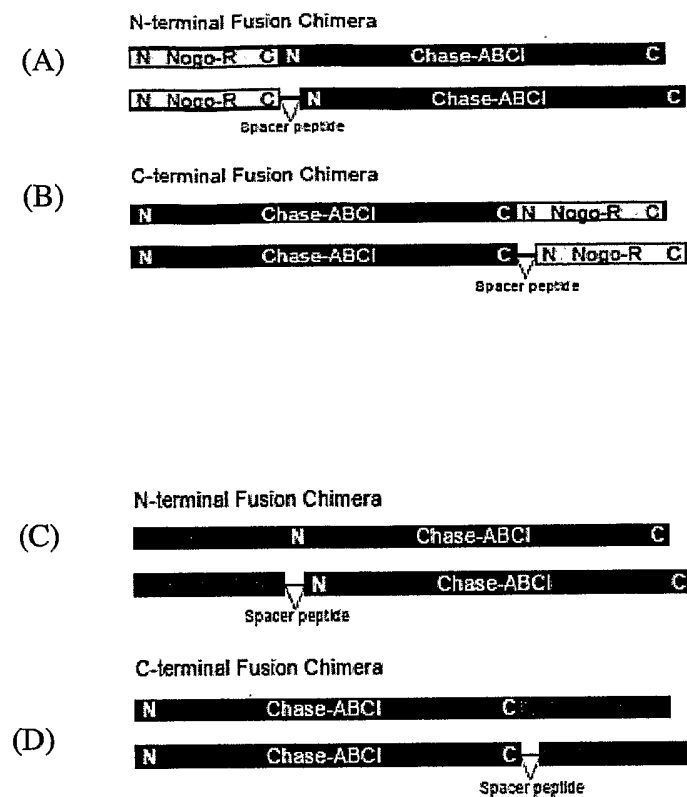


FIG. 4

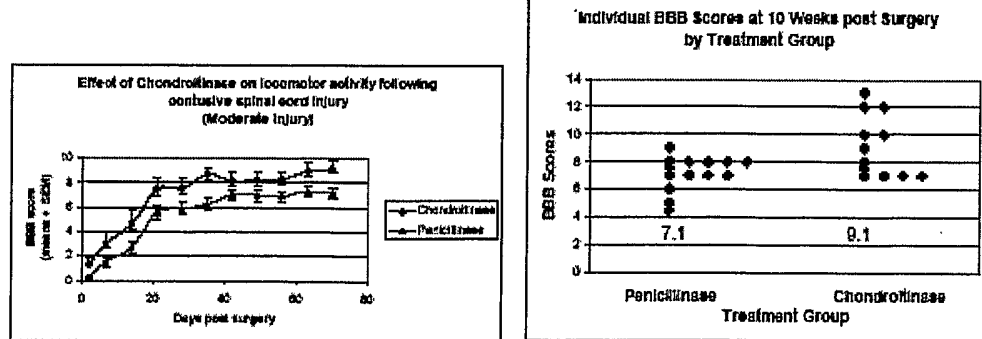
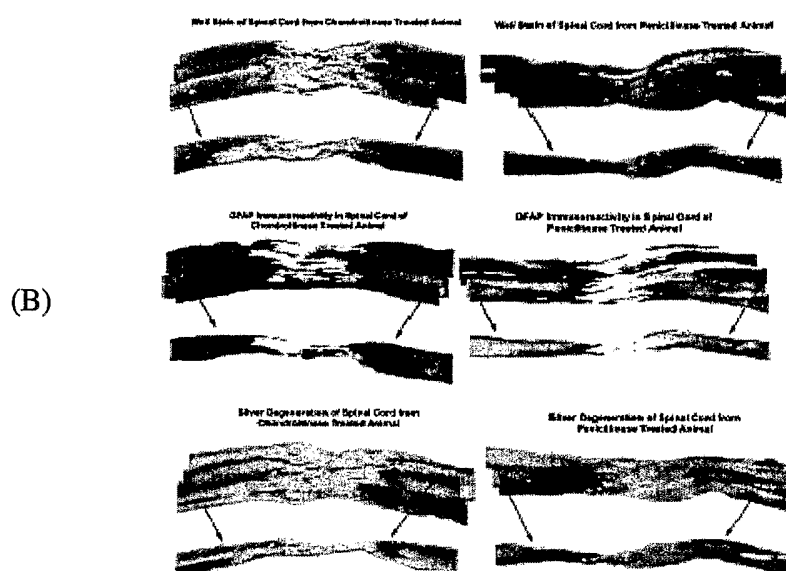
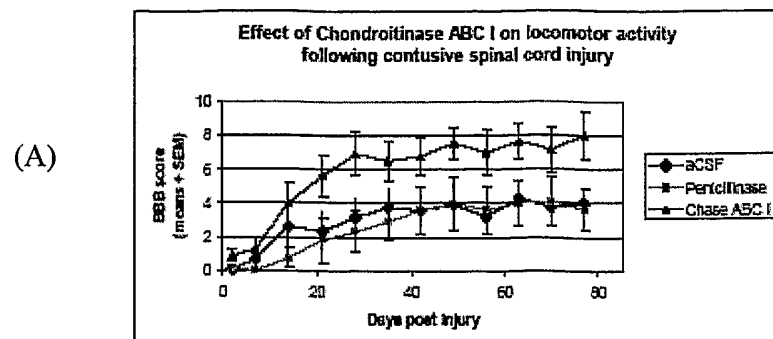


Figure 5

SEQUENCE LISTING**SEQ ID NO: 1** Chondroitinase ABC I protein**ORIGIN**

atsnpa fdpknlmqse iyhfaqnnpl adfssdknsi
61 ltlsdkrsim gnqsllwkwk ggssftlhkk livptdkeas kawgrsstpv fsfwlynekp
121 idgyltidfg eklistseaq agfkvklfdt gwrtvgvsln ndlenremlt natntssdgt
181 qdsigrlsga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
241 hnvkpqlpvt penlaaidli qrlinefvg geketnlale enisklksdf dalnThtlan
301 ggtqgrhlit dkqiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdpktqa
361 qlkqmyllmt khlldqgfvk gsalvtthhw gyssrwwyis tlmsdalke anlqtqvyyds
421 llwysrefks sfdmkvsads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
541 amvsawiyasn pevgiplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
601 tqnestaifg etitpaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr
661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdlrspk phlqmqrger
721 gfsqtsleg qygmmafnli ypanlerfdp nftakksvla adnhlfigs ninssdknkn
781 vettlfqhai tptlntlwq gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
841 hqvsauenkrn qptegnfssa widhstrpkd asyeymvfld atpekmrema qkfrenngly
901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtlivsavtp
961 dlnmtrqkaa tpvtinvtn gkwqsadkns evkyqvsgdn teltftsyfg ipqeiklsp
1021 p

SEQ ID NO: 2 NΔ20 ABCI (A₄₅-N₁₀₂₃), protein

aqnnpl adfssdknsi
61 ltlsdkrsim gnqsllwkwk ggssftlhkk livptdkeas kawgrsstpv fsfwlynekp
121 idgyltidfg eklistseaq agfkvklfdt gwrtvgvsln ndlenremlt natntssdgt
181 qdsigrlsga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
241 hnvkpqlpvt penlaaidli qrlinefvg geketnlale enisklksdf dalnThtlan
301 ggtqgrhlit dkqiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdpktqa
361 qlkqmyllmt khlldqgfvk gsalvtthhw gyssrwwyis tlmsdalke anlqtqvyyds
421 llwysrefks sfdmkvsads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
541 amvsawiyasn pevgiplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
601 tqnestaifg etitpaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr
661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdlrspk phlqmqrger
721 gfsqtsleg qygmmafnli ypanlerfdp nftakksvla adnhlfigs ninssdknkn
781 vettlfqhai tptlntlwq gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
841 hqvsauenkrn qptegnfssa widhstrpkd asyeymvfld atpekmrema qkfrenngly
901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtlivsavtp
961 dlnmtrqkaa tpvtinvtn gkwqsadkns evkyqvsgdn teltftsyfg ipqeiklsp
1021 p

SEQ ID NO: 3 NΔ60 ABCI (F₈₅-N₁₀₂₃) protein

ftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfvkldft gwrtvgvsln ndlenremlt natntsssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
 241 hnvkpqlpvt penlaaidli rqrlinefvg geketnlale enisklksdf dalnhtlan
 301 ggtqgrhlit dkiiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsavtthhw gyssrwwyis tlmsdalke anlqtqvyyds
 421 llwysrefks sfmdmksads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
 601 tqnestaifg etitpaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdn
 661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdlrspk philmqrger
 721 gfsgrssleg qygmmafnl ypanlerfdp nftakksvla adnhlfigs ninssdknkn
 781 vettlfqhai tptlntlwinn gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
 841 hqvsauenknr qptechnfssa widhstrpkd asyeymvfld atpekmgema qkfrenngly
 901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtlavsavtp
 961 dlnmtrqkaa tpvtinvtn gkwqsadkns evkyqvsgdn telftsyfg ipqeiklspl
 1021 p

SEQ ID No. 4: NΔ60 CΔ80 ABCI (F₈₅-A₉₄₂) protein

ftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfvkldft gwrtvgvsln ndlenremlt natntsssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
 241 hnvkpqlpvt penlaaidli rqrlinefvg geketnlale enisklksdf dalnhtlan
 301 ggtqgrhlit dkiiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsavtthhw gyssrwwyis tlmsdalke anlqtqvyyds
 421 llwysrefks sfmdmksads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
 601 tqnestaifg etitpaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdn
 661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdlrspk philmqrger
 721 gfsgrssleg qygmmafnl ypanlerfdp nftakksvla adnhlfigs ninssdknkn
 781 vettlfqhai tptlntlwinn gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
 841 hqvsauenknr qptechnfssa widhstrpkd asyeymvfld atpekmgema qkfrenngly
 901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk pa

SEQ ID NO: 5 Chondroitinase AC protein Locus 1HMW_A**ORIGIN**

1 mkklfvtciv ffsilspall iaqqtgtael imkrvmldlk kplrnmdkva eknlnlqpd
 61 gswkdvpkykd damtnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkkp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypvqfv hyeeglqydy sylqhgpqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrgv srpdilnkka
 301 ekkrllvakm idlkhtewa daiardstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyltd
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydslqakk awffffdkeiv clgaginsna
 481 penittlnq swlengpvist agktgrgkit tfkaqgqfwl lhdaigyyfp eganlslstq

541 sqkgnwfhin nshskdevsg dvfklwinhg arpenaqyay ivlpginkpe eikkyngtap
 601 kvlantnqlq avyhqqldmv qaifytagkl svagieietd kpcavlikhi ngkqviwaad
 661 plqkektavl sirdlktgkt nrvkidfpqq efagatvelk

SEQ ID NO: 6 C Δ 200 AC (Q₂₃-T₅₀₀) protein

qqtgtael imkrvmldlk kplrnmdkva eknlnlqpd
 61 gswkdvpykd damtnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkkp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypvqfv hyeeglqydy sylqhgpqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyltd
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydsqlakk awfffdkeiv clgaginsna
 481 penittlnq swlngpvist

SEQ ID NO: 7 C Δ 220 AC (Q₂₃-A₄₈₀) protein

qqtgtael imkrvmldlk kplrnmdkva eknlnlqpd
 61 gswkdvpykd damtnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkkp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypvqfv hyeeglqydy sylqhgpqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyltd
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydsqlakk awfffdkeiv clgaginsna

SEQ ID NO: 8 N Δ 20 C Δ 200 AC (L₄₃-T₅₀₀) protein

lrnmdkva eknlnlqpd
 61 gswkdvpykd damtnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkkp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypvqfv hyeeglqydy sylqhgpqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyltd
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydsqlakk awfffdkeiv clgaginsna
 481 penittlnq swlngpvist

SEQ ID NO: 9 N Δ 50 C Δ 200 AC (T₇₄-T₅₀₀) protein

tnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkkp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypvqfv hyeeglqydy sylqhgpqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyltd
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydsqlakk awfffdkeiv clgaginsna
 481 penittlnq swlngpvist

SEQ ID NO: 10 NΔ100 CΔ200 AC (S₁₂₃-T₅₀₀) protein

srnwwhne iatpqalgem lilmrygkqp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypqfv hyeeglqydy sylqhgqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrgv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyld
 421 rpltklwgeq gsndfaggvs dgvygasaya ldydsqlakk awffffdkeiv clgaginsna
 481 penittlnq swlulgqvist

SEQ ID NO: 11 NΔ50 CΔ275 AC (T₇₄-L₄₂₆) protein

tnwlppn hllqletiiq ayiekdshyy gddkvfdqis kafkywydsd
 121 pksrnwwhne iatpqalgem lilmrygkqp ldealvhklt ermkrgepek ktganktdia
 181 lhyfyrallt sdeallsfav kelfypqfv hyeeglqydy sylqhgqlq issygavfit
 241 gvlklanyvr dtpyalstek laifskyyrd sylkairgsy mdfnvegrgv srpdilnkka
 301 ekkrllvakm idlkhtewa daiartdstv aagykiepyh hqfwngdyvq hlrpaysfnv
 361 rmvskrtrrs esgnkenllg rylsdgatni qlrgpeyyni mpvwewdkip gitsrdyld
 421 rpltkl

SEQ ID NO: 12 Chondroitinase B Locus Q46079 protein

ORIGIN

1 mkmlnklagy llpimvllnv apclgqvvas netlyqvvke vkpgglvqia dgtvkdvqli
 61 vsnsgksglp itikalnpgk vfftgdakve lrgehlileg iwfkdnrai qawkshgpgl
 121 vaiygsynri tacvfdfde ansayitssl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnpk pgnaggiri gyrrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacyne lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl
 361 derrkeycaa nrlnkfetphq lmlkgnlffk dkpyvypffk ddyfiagkns wtgnvalgve
 421 kgipvnisan rsaykpvkik diqpiegial dlnaliskgi tgkplswdev rpywlkempg
 481 tyaltarlsa draakfkavi krnkeh

SEQ ID NO: 13 NΔ80 Chase B (G₁₀₆-H₅₀₆) protein

gnrai qawkshgpgl
 121 vaiygsynri tacvfdfde ansayitssl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnpk pgnaggiri gyrrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacyne lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl
 361 derrkeycaa nrlnkfetphq lmlkgnlffk dkpyvypffk ddyfiagkns wtgnvalgve
 421 kgipvnisan rsaykpvkik diqpiegial dlnaliskgi tgkplswdev rpywlkempg
 481 tyaltarlsa draakfkavi krnkeh

SEQ ID NO: 14 NΔ120 Chase B (I₁₄₆-H₅₀₆) protein

ittsl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnpk pgnaggiri gyrrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacyne lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl

361 derrkeycaa nrlkfetphq lmlkgnlffk dkpyvypffk ddyfiagkns wtgnvalgve
 421 kgipvnisan rsaykpvkik diqpiegial dlnaliskgi tgkplswdev rpywlkempg
 481 tyaltarlsdraakfkavi krnkeh

SEQ ID NO: 15 CΔ19 Chase B (Q₂₆-L₄₈₈) protein

qvvas netlyqvvke vkpgglvqia dgtvkdvqli
 61 vsnsgksglp itikalnpgk vfftgdakve lrgehlileg iwfkdgndai qawkshgpgl
 121 vaiygsynri tacvfdfde ansayitssl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnppqk pgnaggiri gyyrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacnyfe lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl
 361 derrkeycaa nrlkfetphq lmlkgnlffk dkpyvypffk ddyfiagkns wtgnvalgve
 421 kgipvnisan rsaykpvkik diqpiegial dlnaliskgi tgkplswdev rpywlkempg
 481 tyaltarl

SEQ ID NO: 16 CΔ120 Chase B (Q₂₆-K₃₉₀) protein

qvvas netlyqvvke vkpgglvqia dgtvkdvqli
 61 vsnsgksglp itikalnpgk vfftgdakve lrgehlileg iwfkdgndai qawkshgpgl
 121 vaiygsynri tacvfdfde ansayitssl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnppqk pgnaggiri gyyrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacnyfe lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl
 361 derrkeycaa nrlkfetphq lmlkgnlffk

SEQ ID NO: 17 NΔ120 CΔ120 Chase B (I₁₄₆-K₃₉₀) protein

itssl tedgkvpqhc ridhcsftdk itfdqvinln
 181 ntaraikdgs vggpgmyhrv dhcffsnppqk pgnaggiri gyyrndigrc lvdsnlfmrq
 241 dseaeiitsk sqenvyygnt ylncqgtmnf rhgdhqvain nfyigndqrf gyggmfvwgs
 301 rhviacnyfe lsetiksrgn aalylnpgam asehalafdm liannafinv ngyaihfnpl
 361 derrkeycaa nrlkfetphq lmlkgnlffk

SEQ ID NO: 18 Choindroitinase AC nucleotide locus CHU27583**ORIGIN**

1 atgaagaaat tattttaac ctgtatagtc ttttctcta tttaagtcc tgctctgctt
 61 attgcacagc agaccggta tcgagaactg attatgaagc gggtgatgct ggaccta
 121 aagcctttgc gcaatatgga taagggtggcg gaaaagaacc tgaatacgct gcagcctgac
 181 ggttagctgga aggatgtgcc ttataaagat gatgccatga ccaattgggtt gccaaacaac
 241 cacctgctac aattggaaac tattatacag gcttatattt aaaaagatag tcactattat
 301 ggcgacgata aagtgttgcc ccagatttcc aaagcttttta agtattggta tgacagcgac
 361 ccgaaaagcc gcaactgggt gcacaatgaa attgccactc cgccaggccct tggtaaatg
 421 ctgatctga tgggttacgg taaaaagccg cttgatgaag cattggtaa taaattgacc
 481 gaaagaatga agcggggcga accggagaag aaaacggggg ccaacaaaac agatatcgcc
 541 ctgcattact tttatcggtc tttgttaacg tctgtatggc cttgccttc cttcgccgt
 601 aaagaattgt ttatccgt acagttgtta cactatgagg aaggccgtca atacgattat
 661 tcctacactgc agcacggtcc gcaattacag atatcgagct acggccgtt atttattacc
 721 ggggtactga aacttgccaa ttacgtttagg gataccccctt atgcttaag taccgagaaa
 781 ctggctatat tttcaaaagta ttaccgcgac agttagtctga aagctatccg tggaaagttat

841 atggattta acgtagaagg ccgcggagta agccggccag acattctaaa taaaaaggca
 901 gaaaaaaaaaaga gggtgctggt ggcgaagatg atcgatctta agcatactga agaatggct
 961 gatgcgatag ccaggacaga tagcacagtt gcggccggct ataagattga gccctatcac
 1021 catcagtct ggaatggtga ttatgtgcaa catttaagac ctgcctattc ttttaatgtt
 1081 cgtatggtga gtaagcggac ccgacgcagt gaatccggca ataaagaaaa cctgctggc
 1141 aggtatttat ctgatggggc tactaacata caattgcgcg gaccagaata ctataacatt
 1201 atgccgtat gggaatggga caagattcct ggcataacca gccgtgatta tttaccgac
 1261 agaccttga cgaagcttg gggagagcag gggagcaatg actttgcagg aggggtgtct
 1321 gatgggttat acggggccag tgcctacgc ttggattacg atagcttaca ggcaaagaaaa
 1381 gcctggttct ttttgacaa agagattgtt tgcttgggtt ccggatcaa cagcaatgcc
 1441 cctgaaaaca ttaccactac ccttaaccag agctggtaa atggcccggt tataagtact
 1501 gcaggtaaaa ccggccgggg taaaataaca acgtttaaag cacagggaca gttctggttg
 1561 ttgcacgatg cgattggta ttactttcct gaagggggca accttagtct gagtacccag
 1621 tcgcaaaaag gcaattggtt ccacatcaac aattcacatt caaaagatga agttctggt
 1681 gatgtattta agcttggat caaccatggt gccaggccag aaaatgcgcgca gtatgcctat
 1741 atcggttgc cgggataaaa caagccggaa gaaattaaaa aatataatgg aacggcaccg
 1801 aaagtccttg ccaataccac ccagctgcag gcagtttac atcagcagtt agatatggta
 1861 caggctatct tctatacagc tggaaaatttta agcgtacgg gcatagaaat tggaaacagat
 1921 aagccatgtg cagtctgtat caagcacatc aatggcaagc aggttaatttgc ggctgccc
 1981 ccattgcaaa aagaaaaagac tgcagtgtt agcatcagg attaaaaaac aggaaaaaaca
 2041 aatcggttaa aaattgattt tccgcaacag gaatttgcag gtgcaacggc tgaactgaaa
 2101 tag

//

SEQ ID NO: 19 Chondroitinase AC nucleic acid deletion NΔ50 C Δ275 (a₂₂₀ – t₁₂₇₈)

atgccccatga ccaattggtt gcccaaacaac
 241 cacctgctac aattggaaac tattatacag gcttatatttggaaaaagatag tcactattat
 301 ggcgacgata aagtgttga ccagatttcc aaagcttta agtattggta tgacagcgc
 361 ccgaaaagcc gcaactggtg gcacaatgaa attgcactc cgcaggccct tggtaaaatg
 421 ctgatcctga tgcgttacgg taaaagccg cttgtatggat cattgggtca taaattgacc
 481 gaaagaatga agcggggcga accggagaag aaaacggggg ccaacaaaac agatatgc
 541 ctgcattact tttatcgatg tttgttaacg tctgtatggg cttgccttc cttcgccgta
 601 aaagaattgtt tttatcccgta acagttgttta cactatgagg aaggccctgca atacgattat
 661 tcctacctgc agcacggtcc gcaattacag atatcgagct acggccgtt atttattacc
 721 ggggtactga aacttgccaa ttacgttagg gataccctt atgcttaag taccgagaaa
 781 ctggctatat tttcaaaagta ttaccgcgac agttatctgtt aagctatccg tggaaatgtt
 841 atggattta acgtagaagg ccgcggagta agccggccag acattctaaa taaaaaggca
 901 gaaaaaaaaaaga gggtgctggt ggcgaagatg atcgatctta agcatactga agaatggct
 961 gatgcgatag ccaggacaga tagcacagtt gcggccggct ataagattga gccctatcac
 1021 catcagtct ggaatggtga ttatgtgcaa catttaagac ctgcctattc ttttaatgtt
 1081 cgtatggtga gtaagcggac ccgacgcagt gaatccggca ataaagaaaa cctgctggc
 1141 aggtatttat ctgatggggc tactaacata caattgcgcg gaccagaata ctataacatt
 1201 atgccgtat gggaatggga caagattcct ggcataacca gccgtgatta tttaccgac
 1261 agaccttga cgaagctt

SEQ ID NO: 20 Chondroitinase B nuclei acid Locus CHU27584

ORIGIN

1 atgaagatgc tgaataaaact agccggatac ttattgcgcg tcatgggtct gctgaatgtg
 61 gcaccatgtt taggtcagggt tggcttca aataccaggat tggaaaggag

SEQ ID NO: 21 Chondroitinase B nucleic acid deletion NΔ120 CΔ120 (a₄₃₆ – g₁₁₇₀)

attac tacttcgctt accgaagacg gaaaggtaacc tcaacattgc
481 cgcatacgacc attgcagttt taccgataag atcacttttg accaggtaat taacctgaac
541 aatacagccaa gagctattaa agacggttcg gtgggaggac cggggatgtt ccatcggtt
601 gatcactgtt tttttccaa tccgcaaaaa cgggtaatg ccggaggggg aatcaggatt
661 ggctattacc gtaatgatat aggccgttgt ctggtagact ctaacctgtt tatgcgtcag
721 gattcggaaag cagagatcat caccagcaaa tcgcaggaaa atgtttatttt tgtaataact
781 tacctaattt gccaggcac catgaactt cgtcacggtg atcatcagggtt gcccattaac
841 aattttata taggcaatga ccagcgattt ggatacgggg gaatgtttgt ttggggaaagc
901 aggcattgtca tagcctgtaa ttatTTTgat ctgtccggaaa ccataaaagtc gagggggaaac
961 gcccatttgtt atttaaaccc cgggtctatg gcttcggagc atgctcttgc ttgcataatg
1021 ttgatagccaa caaacgcattt catcaatgtt aatgggtatg ccatccattt taatccattt
1081 gatgagcgcataaaagaata ttgtgcagcc aataggctttaa agttcgaac cccgcaccag
1141 ctaatgttaa aaggcaatct ttcttttaag

SEQ ID NO: 22 Chondroitinase ABCI nucleic acid Locus I29953

ORIGIN

1 ggaattccat cactcaatca ttaaatttag gcacaacgat gggctatcag cgttatgaca
61 aatttaatga aggacgcatt ggittcactg tttagccagcg ttctaaagga gaaaaataat
121 gccgatattt cgittactg cactgcaat gacattgggg ctattatcag cgcctataa
181 cgcgatggca gccaccagca atcctgcatt tgatctaaa aatctgtatgc agtcagaaat
241 ttaccatttt gcacaaaata acccatttagc agacttctca tcagataaaa actcaatact
301 aacgttatct gataaacgta gcattatggg aaaccaatct ctttatgga aatgaaagg

361 tggttagtagc ttactttac ataaaaaaact gattgtcccc accgataaaag aagcatctaa
421 agcatgggga cgctcatcta ccccccgtttt ctcattttgg ctttacaatg aaaaaccgat
481 tcatgtttat cttaactatcg atttcggaga aaaactcatt tcaaccagtg aggctcaggc
541 aggcttaaaa gtaaaattag atttcactgg ctggcgtgtc gtggagatct cttaataaa
601 cgatctgaa aatcgagaga tgaccctaa tgcaaccaat acctccctcg atgtactca
661 agacagcatt gggcgttct taggtctaa agtgcataatg attcgttta aagcgccttc
721 taatgtgagt cagggtgaaa tctataatcg ccgtattatg tttctgtcg atgtgtcg
781 ctaccaatgg tctgattatc aagtaaaaac tcgcttatca gaacctgaaa ttcaatttca
841 caacgtaaag ccacaactac ctgtaaacacc tgaaaattta gcggccattg atcttattcg
901 ccaacgtcta attaatgaat ttgtcgagg tgaaaaagag acaaaccctcg cattagaaga
961 gaatatcagc aaattaaaaaa gtgatttcga tgctcttaat attcacactt tagcaaatgg
1021 tggAACGCAA ggcagacatc tgatcactga taaacaaatc attatttatac aaccagagaa
1081 tcttaactcc caagataaaac aactatttga taattatgtt atttttaggtt attacacgac
1141 attaatgttt aatattagcc gtgcctatgt gctggaaaaaa gatcccacac aaaaggcgca
1201 actaaaggcag atgtacttat taatgacaaa gcatttatta gatcaaggct ttgttaaagg
1261 gagtgcttta gtgacaacccc atcactgggg atacagtct cgtgggtgtt atattccac
1321 gttattatgt tctgtatgcac taaaagaagc gaacctacaa actcaagttt atgattcatt
1381 actgtggat tcaatgtgagt taaaaggtag ttttgcattatg aaagttagtgc ctgatagctc
1441 tgatctatgat tattcaata ccttatctcg ccaacattta gccttattat tactagagcc
1501 tgatgtatca aacgttatca accttagttaa tacitcagc cattatatca ctggcgcatt
1561 aacgcaagtgc ccaccgggtt gtaaagatgg ttacgcctt gatggtagc catggcgcaca
1621 tgaaggcaac tatccgggtt actcttccc agcctttaaa aatgcctctc agcttattta
1681 ttattacgc gatacaccat ttcaatgggg tgaaagggtt tggataacc tgaaaaaaagc
1741 gatggttca gcgtggatct acagtaatcc agaagttgga ttaccgcctt caggaagaca
1801 ccctttaac tcacccatcg taaaatcgt cgcctcaaggc tattactggc ttgcattgtc
1861 tgcaaaatca tcgcctgata aaacacttgc atctattttat cttgcgttta gtgataaaaac
1921 acaaaaatgaa tcaactgctt ttttggaga aactattaca ccagcgtt tacctcaagg
1981 ttctatgcc ttatggcg gtgttttgg tattcatgt tggcaagata aatggtgac
2041 actgaaagct tataacacca atgtttggtc atctgaaatt tataacaaag ataaccgttta
2101 tggccgttac caaagtcatg gtgtcgctca aatagttagt aatggctcgc agcttccaca
2161 gggctatcag caagaagggtt gggattggaa tagaatgcaa gggcaacca ctattcacct
2221 tcctttaaaa gacttagaca gtcctaaacc tcataccctt atgcaacgtg gagagcgtgg
2281 attagcgga acatcatccc ttgaagggtca atatggcatg atggcattcg atcttattta
2341 tcccgccat cttagcggtt ttgatctttaa ttcaatgtcg aaaaagagtg tattagccgc
2401 tgataatcac ttatttttt ttgttagcaa tataaatagt agtgataaaa ataaaaatgt
2461 tgaaacgacc ttatccaaac atgcccattac tccaacatttta aatccctt ggattatgg
2521 acaaaaagata gaaaacatgc cttatcaac aacactcaa caaggtgatt ggttaattga
2581 tagcaatggc aatggttact taattactca agcagaaaaaa gtaaatgtaa gtgcacaaca
2641 tcaggttca gggaaaaata aaaatcgcca accgacagaa gggaaacttta gtcggcatg
2701 gatcgatcac agcactcgcc ccaaagatgc cagttatgatg tattatggctt ttttagatgc
2761 gacacctgaa aaaatggggag agatggcaca aaaattccgt gaaaataatg ggttatatca
2821 ggttcttgcgt aaggataaag acgttcatat tattctcgat aaactcagca atgtaacggg
2881 atatgcctttt tattatggcg catcaattga agacaaatgg atcaaaaagg ttaataaaacc
2941 tgcaattgtg atgactcatc gacaaaaaaga cacttttattt gtcagtgcatg ttacaccgt
3001 tttaaatatg actcgccaaa aagcagcaac tcctgtcacc atcaatgtca cgattaatgg
3061 caaatggcaa tctgcgtataaaaatgtg aatgtaaatat caggtttctg gtgataacac
3121 tgaactgacg ttacgagtt acittggat tccacaagaa atcaaaactct cgcactccc
3181 ttgatttaat caaaaagaacg ctcttgcgtt cttttttat ttgcaggaaa tctgattatg
3241 ctaataaaaaa accctttagc ccacgcgggtt acattaaagcc tctgtttatc attaccgc
3301 caagcattac ccactctgtc tcatgaagct ttccggcgtata ttatctttt tgaagggtgaa

3361 ttacccaata cccttaccac ttcaaataat aatcaattat cgctaagcaa acagcatgct
 3421 aaagatggtg aacaatcaact caaatggcaa tatcaaccac aagcaacattt aacactaaat
 3481 aatattgtta attaccaaga tgataaaaaat acagccacac cactcacttt tatgatgtgg
 3541 atttataatg aaaaacctca atctccccca ttaacgttag cattaaaca aaataataaa
 3601 attgcactaa gtttaatgc tgaacttaat ttacggggg ggcgaggat tgctgttc
 3661 ttctgtata tgcaaggctc tgcgacaggt caacttgatc aattagtgtat caccgctcca
 3721 aaccaagccg gaacactttt tttgtatcaa atcatcatga gtgtacggt agacaatgt
 3781 tggcagttac ctgactatca aacaccctac gtaaataacg cagtaaacac gatgggttagt
 3841 aaaaactgga gtgcattattt gatgtacgt cagatgttc aagcccatta ccctacttta
 3901 aacttcgata ctgaatttcg cgatgaccaa acagaaatgg cttcgattta tcagcgcttt
 3961 gaatattatc aaggaattcc

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SEQ ID NO: 23 TAT fusion chondroitinase ABCI nucleic acid

ggcgtaaaaagcgctgtcaacgtcg
 tgggtgggtgggtgccaccagca atcctgcatt tgatcctaaa aatctgtgc agtcagaaat
 241 ttaccatttt gcacaaaata acccattagc agactctca tcagataaaa actcaatact
 301 aacgttatct gataaacgta gcattatggg aaaccaatct ctttatgga aatggaaagg
 361 tggtagtagc ttactttac ataaaaaaact gattgtcccc accgataaag aagcatctaa
 421 agcatggggc cgctcatcta ccccccgtttt ctcatttgg ctttacaatg aaaaaccgat
 481 tggatgttat ctactatcg atttggaga aaaactcatt tcaaccagtg aggctcaggc
 541 aggcttaaa gtaaaattag atttcactgg ctggcgtgt gtgggaggtct cttaaataaa
 601 cgatctgaa aatcgagaga tgaccttaaa tgcaaccaat acctccctcg atggactca
 661 agacagcatt gggcggtttt taggtgctaa agtgcataatg attcgttta aagcgcccttc
 721 taatgtgagt cagggtgaaa tctatatcga cctgtattatg ttctgtcg atgatgtcg
 781 ctaccaatgg tctgattatc aagtaaaaac tcgcttatca gaacctgaaa ttcaattca
 841 caacgtaaag ccacaactac ctgtAACACC tgaaaattta gcccgttgc atctattcg
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 1201 actaaacgcg atgtacttat taatgacaaa gcatttattt gatcaaggct ttgttaaagg
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 1501 tggatgtca aacgttatca acttagttaa tactttcagc cattataca ctggcgatt
 1561 aacgcacgtg ccaccgggtg gtaaaagatgg ttacggccct gatggatcag catggcgaca
 1621 tggaaacgcaac tatccgggtt actcttccc agcctttaaa aatgcctctc agcttattta
 1681 ttattacgc gatacaccat ttctcgtggg tgaaaagtgg tggataaacc tgaaaaaaagc
 1741 gatggttca gcgtggatct acagtaatcc agaagttgga ttaccgcttg caggaagaca
 1801 cccttttaac tcacccctgt taaaatcagt cgctcaaggc tattactggc ttgcattgtc
 1861 tggcaaaatca tcgcctgata aaacacttgc atctattttat cttcgatgtt gatggaaaac
 1921 aacaaatgaa tcaactgcta ttttggaga aactattaca ccagcgtctt tacctcaagg
 1981 ttctatgcc ttaatggcg gtgcgtttgg tattcatcgat tggcaagata aaatggtgac
 2041 actgaaagct tataacacca atgtttggc atctgaaattt tataacaaag ataaccgtta
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2281 atttagcgg aacatcatccc ttgaaggta atatggcatg atggcattcg atcttattta
 2341 tccgc当地 cttagcgtt ttgatctaa ttcaactcgaaaaagatgt tattagccgc
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 2461 tgaaacgacc ttattccaac atgccattac tccaaacatta aataccctt ggattaatgg
 2521 acaaaagata gaaaacatgc cttatcaaac aacactcaa caaggtgatt ggtaattga
 2581 tagcaatggc aatggttact taattactca agcagaaaaa gttaatgtaa gtcgccaaca
 2641 tcaggttca gcggaaaata aaaatgc当地 accgacagaa ggaaacttta gctggcatg
 2701 gatcgatcac agcactcgcc ccaaagatgc cagttatgat tataatgtct ttttagatgc
 2761 gacacctgaa aaaatggag agatggcaca aaaatccgt gaaaataatg ggtttatatca
 2821 ggttctcgt aaggataaag acgttcatat tattctcgat aaactcagca atgtAACGGG
 2881 atatgc当地 tatcagccag catcaattga agacaaatgg atcaaaaagg ttaataaaacc
 2941 tgcaattgtg atgactcatc gacaaaaaaga cactttattt gtcagtgcatg ttacacctga
 3001 tttaaatatg actgc当地 aagcagcaac tcctgtcacc atcaatgtca cgattaatgg
 3061 caaatggcaa tctgctgata aaaatagtga agtgaardat caggttctg gtgataacac
 3121 tgaactgacg ttacgagtt actttgtat tccacaagaa atcaaactct cgccactccc
 3181 ttgatTTAT caaaaagaacg ctcttgcgtt cctttttat ttgcaggaaa tctgattatg
 3241 ctaataaaaaa accctttagc ccacgc当地 acatTAAGCC tctgtttatc attaccgc当地
 3301 caagcattac ccactctgat tcatgaagct ttccgc当地 tttatcttt tgaagggtgaa
 3361 ttacccaata cccttaccac ttcaaataat aatcaattat cgctaagcaa acagcatgct
 3421 aaagatggtg aacaatcaat caaatggcaa tatcaaccac aagcaacattt aacactaaat
 3481 aatattgttta attaccaaga tgataaaaaat acagccacac cactcactt tatgtatgtgg
 3541 atttataatg aaaaacctca atcttccccca ttaacggtt catttaaca aataataaaa
 3601 attgcactaa gtttaatgc tgaacttaat ttacgggtt ggc当地 ggtt tgc当地 tcc
 3661 ttccgtgata tgcaaggctc tgc当地 gaggtt caacttgatc aatttgatgat caccgcttca
 3721 aaccaagccg gaacactctt ttgtatcaa atcatcatga gtgtaccgtt agacaatgt
 3781 tggc当地 ctgactatca aacacccttac gtaataacg cagtaaacac gatggtttagt
 3841 aaaaactgga gtgc当地 tattt gatgtacgat cagatgttca aagccattt ccctacttta
 3901 aacttc当地 gtaatttcg cgtatgaccaa acagaaatgg ctgc当地 ttagcgtt
 3961 gaatattatc aaggaattcc

SEQ ID NO: 24 An HIV TAT sequence and Gly penta linker protein

G R K K R R Q R R R G G G G G

SEQ ID NO: 25 Tat sequence and Gly penta linker nucleic acid

ggt cgt aaa aag cgt cgt caa cgt cgt ggt ggt ggt ggt

SEQ ID NO: 26 Present Invention Chondroitinase ABC II Nucleic acid

ttaccactctgtctcatgaagcttc当地ggc当地gatatttatcttttgaaggc当地gatattccaaatc当地cttaccacttcaaaataataatcaattatcg
 ctaagcaacagcatgctaaagatggtaacaatcactcaaatggcaatcatcaaccacaagcaacattaacactaaataataatgttaatt
 accaagatgataaaaatcagccacaccactcactttatgatgtggatttataatgaaaaacctcaatcttccc当地ttaacgttagtatttca
 aacaaaataataaaatgc当地actaagtttatgctgacttaattttacggggtggc当地gaggatttgc当地tcc当地tgc当地gatatgc当地aggctc
 tgc当地gaggctcaacttgc当地atttgc当地acttgc当地acccttgc当地ttaacacactcacttgc当地acttgc当地gaggctt
 gacaatcgtggc当地gaggcttgc当地acttgc当地acccttgc当地ttaacacactcacttgc当地acttgc当地gaggctt
 atgtacgatcagatgttcaagccc当地taccctactttaaacttc当地gataacttgc当地gatgaccaaacagaaatggcttgc当地atttgc当地
 cgcttgc当地atttgc当地aggatcgttagtataaaaaattactcc当地gatatgtctgacttgc当地acttgc当地gatgaccaaacagaaatggcttgc当地atttgc当地
 acacaacacgctgatggctcaatcacaaggaaaagcccttgc当地acccttgc当地accggcaacattttatgaaagtc当地gaggatgtt
 gggactcaaaaagc当地atttgc当地acttgc当地acttgc当地gatgaccaaacgcttgc当地acttgc当地tatttgc当地acttgc当地gattca
 ttatc当地gacttgc当地tagaaaaatttgc当地acttgc当地tatttgc当地acttgc当地gatgaccaaacagaaatggcttgc当地atttgc当地

tcaaatttactcatgttggttaccaaaccagagaactttatgcgttgcattggccgtatgttgcacaaaataacccttagcccc
 ccactcaacaagctatgttgcacacgcccacaggacgttttggaaaagataatgaaattgttgcataatgtcgatatttcataat
 actcaatttgcataatggatgataaaaagcttattgtatgcgttgcattcaacaacgtcaacaacgccttagcgcactgcaagtgctaa
 aataaaaaccattctaagctcaaaaggtgttgcgttgcgttgcattctatgttgcattcaacacattaccccgttatgcata
 aagatgcattttgggttagcaccctgttgcattaaatgttgcatttgcgttgcattctacttcgcacatgagcattaaagatgtt
 ttgttaaaaatgcggatctacaccaaagagacacaaattcgttgcattaaatgttgcgttgcattcaactgggtgcataaaataggatgc
 cggcatttaaatggatggcattagcaggaaccccagatggcaacaaaagttgataccacattatccgcgcgttatgcata
 aacaaaacgcatttgcaggcattaaacgcgttgcatttgcgttgcatttgcattcaatggcaatacaacgaa
 gagcatcgacccaatcaccacaaacaaagctggctgcgcattagcgcgcgttgcatttgcgttgcattcaatggcaatacaacgaa
 acaaccgttatggcgttataatggacaatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 ggaatagatattccaggtaacaaactattcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 aatgttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 acaacaaaagcttaagggcaatataatccatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 acgaccgaaacaaacactattccaggtaatggcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 aactttaataatgcagataatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 acatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 atgaatatgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 ataaaataacccaaagaagaggatgtttttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 tatgttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 gatgataaaggtaatcaaatcgaagttgttattctgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 aggaatatggaaattaaacgacacctaaccgggttatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt
 aacacctactgttatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatttgcatt

SEQ ID NO: 27 Present Invention Chondroitinase ABC II protein

>_ ABC (present invention) 990 aa vs.
 >_ ABC (mature) 990 aa

scoring matrix: , gap penalties: -12/-2
 98.3% identity; Global alignment score: 6393

10	20	30	40	50	60
457676 LPTLSHEAFGDIYLFE GELPNIL TTSNNNQLSLSKQHAKDGEQSLKWQYQPQATLTLNNI
- LPTLSHEAFGDIYLFE GELPNIL TTSNNNQLSLSKQHAKDGEQSLKWQYQPQATLTLNNI
10	20	30	40	50	60

70	80	90	100	110	120
457676 VNYQDDKNTATPLTFMMWIYNEKPQSSPLTLAFKQNNKIALSFNAELNFTGWRGIAVPFR
- VNYQDDKNTATPLTFMMWIYNEKPQSSPLTLAFKQNNKIALSFNAELNFTGWRGIAVPFR
70	80	90	100	110	120

130	140	150	160	170	180
457676 DMQGSATGQLDQLVITAPNQAGTLFFDQIIMSVPLDNRWAVPDYQT PYVNNAVNTMVSKN
- DMQGSVTGQLDQLVITAPNQAGTLFFDQIIMSVPLDNRWAVPDYQT PYVNNAVNTMVSKN
130	140	150	160	170	180

190	200	210	220	230	240
457676 WSALLMYDQM FQAHYPTLNFDTEFRDDQTEMAS IYQRFEYYQGIRSDKKITPDMLDKHLA
- WSALLMYDQM FQAHYPTLNFDTEFRDDQTEMAS IYQRFEYYQGIRSDKKITPDMLDKHLA
190	200	210	220	230	240

— WSALLMYDQMFQAHYPTLNFDTEFRDDQTEMASRYQRFEYYQGIRSDKKITPDMLDKHLA
 — 190 200 210 220 230 240
 — 250 260 270 280 290 300
 457676 LWEKLGLTQHADGSITGKALDHPNRQHFMKVEGVFSEGTQKALLDANMLRDVGKTLQTA
 — :::
 — LWEKLVLTQHADGSITGKALDHPNRQHFMKVEGVFSEGTQKALLDANMLRDVGKTLQTA
 — 250 260 270 280 290 300
 — 310 320 330 340 350 360
 457676 IYLRSDSLSATGRKKLEERYLLGTRYVLEQGFTRGSGYQIITHVGYQTRELFDAWFIGRH
 — :::
 — IYLRSDSLSATDRKKLEERYLLGTRYVLEQGFHRGSGYQIISHVGYQTRELFDAWFIGRH
 — 310 320 330 340 350 360
 — 370 380 390 400 410 420
 457676 VLAKNNLLAPQQAMMWNATGRIFEKDNEIVDANVDILNTQLQWMIKSLLMLPDYQQRQ
 — :::
 — VLAKNNLLAPQQAMMWNATGRIFEKNNEIVDANVDILNTQLQWMIKSLLMLPDYQQRQ
 — 370 380 390 400 410 420
 — 430 440 450 460 470 480
 457676 QALAQLQSWLNKTILSSKGVAGGFSDGSIFHHSQHYPAYAKDAFGLAPSVDSPF
 — :::
 — QALAQLQRWLNLKTILSSKGVAGGFSDGSIFHHSQHYPAYAKDAFGLAPSVDSPF
 — 430 440 450 460 470 480
 — 490 500 510 520 530 540
 457676 RLSTSAHEHLKDVLCKMRYTKEQIPVVLSGRHPTGLHKIGIAPFKWMALAGTPDGKQK
 — :::
 — RLSTSAHERLKDVLLKMRYTKEQIPAVLSGRHPTGLHKIGIAPFKWMALAGTPDGKQK
 — 490 500 510 520 530 540
 — 550 560 570 580 590 600
 457676 LDTLSAAYANLDNKTHFEGINAESPVGAWAMNYASMAIQRRASTQSPQQSWLAIARGF
 — :::
 — LDTLSAAYAKLDNKTHFEGIKAESPVGAWAMNYASMAIQRRASTQSPQQSWLAIARGF
 — 550 560 570 580 590 600
 — 610 620 630 640 650 660
 457676 SRYLGNESYENNRYGRYLQYGQLEIIIPADLTQSGFSHAGWDWNRYPGTTTIHLPYNEL
 — :::
 — SRYLGNESYENNRYGRYLQYGQLEIIIPADLTQSGFSHAGWDWNRYPGTTTIHLPYNEL
 — 610 620 630 640 650 660
 — 670 680 690 700 710 720
 457676 EAKLNQLPAAGIEEMLLSTESYSGANTLNNNSMFAMKLHGSKYQQSLRANKSYFLFDN
 — :::
 — EAKLNQLPAAGIEEMLLSTESYSGANTLNNNSMFAMKLHGSKYQQSLRANKSYFLFDN
 — 670 680 690 700 710 720
 — 730 740 750 760 770 780
 457676 RVIALGSGIENDDKQHTTETTLFQFAVPKLQSVIINGKVNQLDTQLTLNNADTLIDPAG
 — :::
 — RVIALGSGIENDDKQHTTETTLFQFAVPKLQSVIINGKVNQLDTQLTLNNADTLIDPAG
 — 730 740 750 760 770 780
 — 790 800 810 820 830 840
 457676 NLYKLTKGQTVKF SYQKQHSLDDRNSKPTEQLFATAVISHGKAPSNENEYAIAIEAQNN
 — :::
 — NLYKLTKGQTVKF SYQKQHSLDDRNSKPTEQLFATAVISHGKAPSNENEYAIAIEAQNN

	790	800	810	820	830	840
	850	860	870	880	890	900
457676	KAPKYTVLQHNDQLHAVKDKITQE E GY G FFEATKLKSADATLLSSDAPVMVMAKIQNQQL ::::::::::: ::::::::::::::::::::: ::::::::::::::::::::: :::::::::::::::::::::					
-	KAPEYTVLQHNDQ P HAVKDKITQE E GY A FFEATKLKSADATLLSSDAPVMVMAKIQNQQL 850	860	870	880	890	900
	910	920	930	940	950	960
457676	TLSIVNPDLNL Y QGREKDQFDKG N QIEVS V YSRHWLTAESQSTNSTITVKGIWKLTT P Q ::::::::::: ::::::::::::::::::::: ::::::::::::::::::::: :::::::::::::::::::::					
-	TLSIVNPDLNL Y QGREKDQFDKG N QIEVS V YSRHWLTAESQSTNSTITVKGIWKLTT P Q 910	920	930	940	950	960
	970	980	990			
457676	PGVI I KHHNNNTLITTTTIQATPTVINLVK ::::::::::: :::::::::::::::::::::					
-	PGVI I KHHNNNTLITTTTIQATPTVINLVK 970	980	990			

SEQ ID NO: 28 Present Invention Chondroitinase ABC I nucleic acid

gccaccagcaatcctgcatttgcataaaaaactgtatgcaggcagtcagaattttgcacaaaataacccattagcagacttcata
agataaaaaactcaataactaacgttatctgatcaaactgtagcattatggaaaccaatctttatggaaatggaaaggtagtttt
cittacataaaaaactgattgtccccaccgataaagaagcatctaaagcatgggacgctatccaccccggttctcatggcttac
aatgaaaaaccgattgtatggttatcttactatcgatccggagaaaaactcattcaaccagtggctcaggcaggcttaaagtaaatt
agatttcactggctgggtactgtggagtcattaaataacgatctgaaaatcgagagatgaccctaaatgcaaccaataccctctg
atggactcaagacagcattggcgtttagtgctaaagtgcatagtattcgtttaaagcgcctctaattgtgagtcaggtaaatt
tatatcgaccgtattatgttctgtcgatgtgctgctaccatggctgattatcaagtaaaaactcgcattcagaacctgaaattcaatt
tcacaacgtaaagccacaactaccgttaacaccgtaaaatttagcggccattgatcttattcgcacacgtctaaatgaatttgcggag
gtgaaaaagagacaaacctcgattagaagagaatatcagcaaaattaaaagtgttgcattactcacacttttagcaaaatg
gtggAACGCAAGGCAGACATCTGATCAGTAAACACCTGTTAATTTCAGGAGAGATCTAACTCTCAAGATAAACACAACATTGT
AATTATGTTATTTCAGGTAATTACACGACATTAATGTTAATTAGCCGTGTTATGTGCTGGAAAAGATCCCACACAAAAGGGCGCAACT
AAGCAGATGTACTTATTAAATGACAAGCATTAGATCAAGGCTTGTTAAAGGGAGTGCTTAGTGACNACCCATCAGTGGGATACA
GTTCTCGTTGGTGTATTTCCACGTTATTAAATGTCGATGCACCTAAAGAGCAGACCTACAAACTCAAGTTATGATTCTGATTACTGT
TCACGTTGAGTTAAAAGTAGTTGATGAAAGTAGTGTGATGCTGATCTGATTCTGATTATTCACACCTTATCTGCCAACATTGACCTT
ATTACTAGAGCCTGATGATCAGGAAAGCGTATCAACTTAGTTAATTCTGAGCCATTATCAGTGGCGCATTACGCAAGTGCACCG
GTGGTAAAGATGGTTACGCCCTGATGGTACAGCATGGCGACATGAAGGCAACTATCCGGCTACTTTCCAGGCTTAAAAGCCT
CTCAGCTTATTATTACGCGATACACCATTCTGAGTGGGTTAAAGGTGGTGAATAGCCTGAAAAAGCGATGGTTAGCGTGGATC
ACAGTAATCCAGAAGTGGATTACCGCTTGAGGAAGACACCCCTTAACTCACCTCGTTAAATCAGTCGCTCAAGGCTTACTGGCT
TGCCATGTCGAAACATCGCCTGATAAAACACTGCACTATTATCTGCGATTAGTGATAAAACACAAAATGAATCAACTGCTT
GGAGAAACTATTACACCGCTTACCTCAAGGGTCTATGCCCTTAATGGCGGTGCTTGGTATTCTGTTGGCAAGATAAAATGGT
ACACTGAAAGCTTAAACACCAATGTTGGTCACTGTTAAACAAAGATAACCGTTATGCCGTACCCAGGCTTAAAGTCATGGTGC
CAATAAGTGAGTAATGGCTCGAGCTTCACAGGGCTATCAGCAAGAAGGGTGGATTGGAAATAGTGCCAGGGCAACCATC
CTCCCTTAAGACTAGACAGCTCAAACCTCATACCTTAATGCAACGTGGAGAGCGTGGATTAGCGGAACATCATCCCTGAAGG
CTAATATGGCATGATGGCATTGCTGATCTTATTATCCGCCAATCTGAGCGTTGATCTAATTCTGCGAAAAAGAGTGATTAG
GCCGCTGATAATCACTTAATTGTTATTGGTAGCAATAATAGTAGTGATAAAATAAAATGGTAAACGACCTTATCCAACATGCC
ATTAAACCTTGGATTATGGACAAAAGAGATAAAACATGCCATTCAAAACACACTCAACAAGGTGATTGGTTAATTGATAG
GCAATGGTACTTAATTACTCAAGCAGAAAAGTAAAGTAAAGTCGCGCAACATCAGGTTCTGCGGAAAATAAAATGCC
GAAGGAAACTTAGCTGGCATGGATCAGCAGCAGCTGCCAAAGATGCCAGTTAGTGTTATGGCTTITAGATGCGACAC
GAAAATAAGGGAGAGATGGCACAAGGAAATTCCGTGAAATAATGGTTATCAGGGTCTGTAAGGATAAAGACGTC
ATAAAACTCAGCAATGTAACGGGATATGCCATTATGCCAGCAGCATCAATTGAGACAAATGGTAAACCTGCAATTGT
GATGACTCATGACAAGACACTTATTGTCAGTCAGTTACACCTGATTAAATGACTCGCCAAAAGCAGCAACTCTGTC
ACCC

atcaatgtcacgattaatggcaa atggcaatctgctgataaaaaatgtgaagt gaaatcagg ttcgg t gataacactgaactgacgt
ttacagttacttggta ttccacaagaatcaaactctgccactccctga

SEQ ID NO: 29 Present Invention Chondroitinase ABC I protein

365019 ATSNPAFDPKNLMQSEIYHFAQNNPLADFSSDKNSILTSKRSIMGNQSLWKWKGSS

.....
 365019 FTLHKKLIVPTDKEASKAWGRSSTPVFSFWLYNEKPIDGYLTIDFGEKLISTSEAQAGFK
 365019 VKLDFTGWRTVGVLNNNDLENREMTLNATNTSSDGTQDSIGRSLGAKVDSIRFKAPSNSV
 365019 QGEIYIDRIMFSVDDARYQWSDYQVKTRLSEPEIQFHNVKPQLPVTPENLAAIDLIRQRL
 365019 INEFVGGEKETNLALEENISKLKSDFDALNTHTLANGGTQGRHLITDKQIIYQOPENLNS
 365019 QDKQLFDNYVILGNYTTLMFNISRAYVLEKDPTQKAQLKQMYLLMTKHLLDQGFVKGSAL
 365019 VTTHHWGYSSRWWYISTLLMSDALKEANLQTQVYDSLLWYSREFKSSFDMKVSADSSDLD
 365019 YFNTLSRQHLALLLEPDDQKRINLVNTFSHYITGALTQVPPGGKDGLRPDGTAWRHEGN
 365019 YPGYSFPFAFKNASQLIYLLRDTPFSVGESGWNSLKKAMVSAWIYSNPEVGLPLAGRHPLN
 365019 SPSLKSVAQGYYWLAMSAKSSPDKTLASIYLAISDKTQNESTAIFGETITPASLPQGFYA
 365019 FNNGAFGIHRWQDKMVTLKAYNTNWSSEIYNKDNRYGRYQSHGVAQIVSNGSQLSQGYQ
 365019 QEGWDWNRMPGATTIHLPLKDLDSPKPHTLMQRGERGFSGTSSLEGQYGMMAFDLIYPAN
 365019 LERFDPNFTAKKSvlaadnhlifigsniNssdknknvettlfqhaiptlntlwingqki
 365019 ENMPYQTTLQQGDWLIDSNGNGYLITQAEKVNVRQHQVSAENKNRQPTEGNFSSAWIDH
 365019 STRPKDASYEYMVFLDATPEKMGEAQKFRENNGLYQVLRKD KVHIIDKLSNVTGYAF
 365019 YQPASIEDKWIKKVNKP AIVMTHRQKDTLIVSAVTPDLMTRQKAATPV TINVTINGKWQ
 365019 SADKNSEVKYQVSGDNTELFTSYFGIPQEIKLSPLP

hyaluronidase 1 protein, **SEQ ID NO: 30**, Locus HSU96078

hyaluronidase 2 protein, **SEQ ID NO: 31**,

hyaluronidase 3 protein, **SEQ ID NO: 32**, Locus BC012892

hyaluronidase 4 protein, **SEQ ID NO: 33**, AF009010

PH-20 protein, **SEQ ID NO: 34**,

Tat peptide **SEQ ID NO: 35**

ABCI a site-specific mutant designated H501a and Y508a **SEQ ID NO: 36**

SEQ ID NO: 37 Chondroitinase ABCI protein Locus P59807

ORIGIN

1 mpifrftala mtlgllsapy namaatsnpa fdpknlmqse iyhfaqnnpl adfssdknsi
 61 ltsdkrsim gnqslwkwk ggssftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfkvkldft gwravgvsln ndlenreml natntssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvktl sepeiqf

241 hnvkpqlpv penlaaidli qrlinefv gcketnlale enisklksdf dalmihtlan
 301 ggtqgrhlit dkqiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsalvtthhw gyssrwwyis tlmsdalke anlqtqvyyds
 421 llwysrefks sfdmkvsads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
 601 tqnestaifg etippaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr
 661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdldspk philmqrger
 721 gfsgrssleg qygmmafnli ypanlerfdp nftakksvla adnhlifigs ninssdknkn
 781 vettlfqhai tptlntlwq gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
 841 hqvsakenr qptegnfssa widhstrpkd asyeymvfld atpekmgema qkfrenngly
 901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtliavsavtp
 961 dlnmtrqkaa tpvtinvtn gkwqsadkns evkyqvsgdn telftsyfg ipqeiklspl
 1021 pXX

SEQ ID NO: 38 NΔ20 ABCI (A₄₅-N₁₀₂₃), protein

aqnnpl adfssdknsi
 61 ltlsdkrsim gnqslwkqk ggssftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfkvklfdt gwrvagsln ndlenremlt natntssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
 241 hnvkpqlpv penlaaidli qrlinefv gcketnlale enisklksdf dalmihtlan
 301 ggtqgrhlit dkqiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsalvtthhw gyssrwwyis tlmsdalke anlqtqvyyds
 421 llwysrefks sfdmkvsads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
 601 tqnestaifg etippaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr
 661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdldspk philmqrger
 721 gfsgrssleg qygmmafnli ypanlerfdp nftakksvla adnhlifigs ninssdknkn
 781 vettlfqhai tptlntlwq gqkienmpyq ttlqqgdwli dsngngylit qaekvnvsrq
 841 hqvsakenr qptegnfssa widhstrpkd asyeymvfld atpekmgema qkfrenngly
 901 qvlrkdkdvh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtliavsavtp
 961 dlnmtrqkaa tpvtinvtn gkwqsadkns evkyqvsgdn telftsyfg ipqeiklspl
 1021 pXX

SEQ ID NO: 39 NΔ60 ABCI (F₈₅-N₁₀₂₃) protein

ftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfkvklfdt gwrvagsln ndlenremlt natntssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
 241 hnvkpqlpv penlaaidli qrlinefv gcketnlale enisklksdf dalmihtlan
 301 ggtqgrhlit dkqiiyqpe nlnsqdkqlf dnyvilgnyt tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsalvtthhw gyssrwwyis tlmsdalke anlqtqvyyds
 421 llwysrefks sfdmkvsads sdldyfntls rqhlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlkk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdktl asiylaisdk
 601 tqnestaifg etippaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr

661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdldspk phtlmqrger
 721 gfsgtssleg qygmmafnl ypanlerfdp nftakksvla adnhlifigs ninssdknkn
 781 vettlfqhai tptlntlwinn gqkienmpyq ttlqqgdwli dsngngylit qaeckvnvsrq
 841 hqvsauenkr qptegnfssa widhstrpkd asyeymvfld atpekmrema qkfrenngly
 901 qvrlrkdkdyh iildklsnvt gyafyqpasi edkwikkvnk paivmthrqk dtlivsavtp
 961 dlnmtrqkaa tpvtinvtin gkwqsadkns evkyqvsgdn telftsyfg ipqeiklspl
 1021 pXX

SEQ ID NO. 40: NΔ60 CΔ80 ABCI (F₈₅-A₉₄₂) protein

ftlhkk livptdkeas kawgrsstpv fsfwlynekp
 121 idgyltidfg eklistseaq agfkvklidft gwrvavgsln ndlenremlt natntssdgt
 181 qdsigrslga kvdsirfkap snvsqgeiyi drimfsvdda ryqwsdyqvk trlsepeiqf
 241 hnvkpqlpvt penlaaidli rqrlinefvq geketnlale enisklksdf dalnihtlan
 301 ggtqgrhlit dkqiiiyqpe nlnsqdkqlf dnyvilgny tlmfnisray vlekdptqka
 361 qlkqmyllmt khlldqgfvk gsalvttthhw gyssrwwyis tillmsdalke anlqtqvyds
 421 llwysrefks sfdmkvsads sdldyfntls rqlallle pddqkrinlv ntfshyitga
 481 ltqvppggkd glrpdtawr hegnypgysf pafknasqli yllrdtpfsv gesgwnnlk
 541 amvsawiysn pevglplagr hpfnspslks vaqgyywlam saksspdkl asiyaisdk
 601 tqnestaifg etitpaslpq gfyafnggaf gihrwqdkmv tlkayntnvw sseiynkdnr
 661 ygryqshgva qivsngsqls qgyqqegwdw nrmegattih lplkdldspk phtlmqrger
 721 gfsgtssleg qygmmafnl ypanlerfdp nftakksvla adnhlifigs ninssdknkn
 781 vettlfqhai tptlntlwinn gqkienmpyq ttlqqgdwli dsngngylit qaeckvnvsrq
 841 hqvsauenkr qptegnfssa widhstrpkd asyeymvfld atpekmrema qkfrenngly
 901 qvrlrkdkdyh iildklsnvt gyafyqpasi edkwikkvnk pa

SEQ ID NO. 41: I. Nucleotide sequence for TAT-ABCI-nΔ20:

1 ggtc gtaaaaagcg tcgtcaacgt cgctgcctc ctaatgcgc acaaaataac
 61 ccattagcag acttctcatc agataaaaaac tcaataactaa cgttatctga taaacgttagc
 121 attatggaa accaatctct ttatggaaa tggaaagggtg gtagtagctt tactttacat
 181 aaaaaactga ttgtccccac cgataaagaa gcatctaaag catggggacg ctcatccacc
 241 cccgtttct cattttggct ttacaatgaa aaaccgattt atggttatot tactatcgat
 301 ttccggagaaaa aactcatttc aaccagttag gctcaggcag gctttaagt aaaatttagat
 361 ttcaactggct ggcgtactgt gggagctct ttaaataacg atcttggaaa tcgagagatg
 421 accttaaatg caaccaatac ctccctgtat ggtactcaag acagcattgg gcttcttta
 481 ggtgctaaag tcgatagttat tcgtttaaa gcgcctcta atgtgagtca gggtgaaatc
 541 tatatcgacc gtattatgtt ttctgtcgat gatgctcgct accaatggc tgattatcaa
 601 gtaaaaactc gcttattcaga acctgaaatt caatttcaca acgtaaagcc acaactacct
 661 gtaacacctg aaaaatttagc ggccattgat ctattcgcc aacgtctaat taatgaattt
 721 gtcggaggtg aaaaagagac aaacctcgca ttagaagaga atatcagcaa attaaaaagt
 781 gatttcgatg ctcttaatac tcacactta gcaaattggtg gaacgcagg cagacatctg
 841 atcaactgata aacaaatcat tatttatcaa ccagagaatc ttaactctca agataaacaa
 901 ctatttgata attatgttat tttaggtaat tacacgacat taatgtttaa tattagccgt
 961 gcttatgtgc tggaaaaaga tcccacacaa aaggcgcaac taaagcagat gtacttatta
 1021 atgacaaagc atttattaga tcaaggctt gttaaaggga gtgccttagt gacnacccat
 1081 cactggggat acagttctcg ttggtggtat atttccacgt tattaatgctc tgatgcacta
 1141 aaagaagcga acctacaaac tcaagtttat gattcattac tgggtattc acgtgagtt
 1201 aaaagttagtt ttgatatgaa agtaagtgc gatagctctg atctagatta tttcaataacc

SEQ ID NO. 42: II. Amino acid sequence for TAT-ABCI-nA20:

grkkrrqrrppqcaqnnpladfssdknsiltlSDKrsimgnqsslwkwkggssflhkklivptdkeaskawgrsstpvfsfwlynekpidgyltdfgeklistseaqagfkvklfdgtgwrtvgvslnndlenremtnatntssdgtqdsigrslgavdssirfkapsnvsqgeiyidrimfsvddaryqwsdyqvktlsepeiqfhnvkpqlpvtpenlaaidlirqrlinefvgeketnlaleenisklksdfdalnhttlanggtqgrhltdkqiiyqpenlnsqdkqlfdnyvilgnytllmfniisrayvlekdptqkaqlkqmyllmtkhlldqgfvkgsalvttihwggyssrwwyistllmsdalkeanlqtqvdydsllwysrefkssfdmkvsadssldyfntlsrqhlllepdqdkirinlvntfshyitgaltqvppggkdgldprdtawrhegnypgysfpafkknasqliyllrdtpfsvgesgwnsllkamvsawiysspnevglplagrhplnspsllksvaqgyywlamssaksspdktlasiylaisdktqnestaifgetitpaslpqgyafnggafgihrwqdkmvtlkayntvwsseiynkdnrygryqshgvaqivsngsqlsrgyqqegwdwnrmpgattihplkldspkphilmqrgergfsgtsslegqyggmmafdliypanlerfdpnftakksvlaadnhlfifgsnninssdknknvettlfqhaitptlntlwingqkienmypyqtlqqgdwlidsngngylitqaekvnvsrqhqvsaenknrqptegnfssawidhstrpkdasyeymvldatpekmgemaqkfrennglyqvrlrkdkdvhilidklsntvgyafyqpasiedkwikkvnkpaivmthrqkdtlivilsavtpdlnmtrqkaatpvtinvtingkwqsadknsevkyqvsgndtelftsyticpoeiklsp

SEQ ID NO. 43: II. Nucleotide sequence for TAT-ABCI-nΔ60:

SEQ ID NO. 44: IV. Amino acid sequence for TAT-ABCI-nΔ60

grkkrrqrrppqcflhkklivptdkeaskawgrsstpvfsfwlynekpidgyltidfgeklistseaqagfkvklfdftgwrtvgvsl
nnrlenremlnatntssdgtdqsigrsgakvdsvirfkapsnvsqgeiyidrimfsddaryqwsdyqvktlsepeiqfhnvkp
qlpvtpenlaaidlirlqrlinefvgeketnlaleenisklksdfdalnthalanggtqgrhltdkqijyqpenlnsqdkqlfdnyvil
gnytllmfnisrayvlekdpqkaqlkqmyllmtkhlldqgfvkgsalvtthhwgyssrwwyistllmsdalkeanlqtqvysdlsll
wysrefkssfmdmkvsadssldyfntlsrqhlllepddqkrinlvntfshyitgaltqvppggkdglrpdtawrhegnypgys
fpafknasqliyllrdtpfsvgesgwnsllkamvsawiysnpevglplagrhlspslksvaqgyywlamaksppdktlasiy
laisdktqnestaifgetitpaslpqgyafnggafgihrwqdkmvtlkayntnvwssieynkdnrygryqshgvaqivsngsqls
qgyqqegwdwnrmpgattihplkldspkphitmqrgergfsgtsslegqygmmafdliypanlerfdpnftakksvlaadn
hlifigsninssdknkvnvetlqhaiptlntlwingqkienmpyqtlqqgdwlidsngngylitqaekvnvsrqhvsaeiknr
qptegnfssawidhstrpkdasyeymvfdatpekmgemaqkfrennglyqvrlkdkdvhiiidklsnvtgyafyqpasiedk
wikkvnkpaivmthrqkdtlivosavtpdlnmtrqkaatpvtinvtingkwqsadknsevkyqvsgdntelttsyfgipqeiklsp
lp

SEQ ID NO. 45: V. Nucleotide sequence for ABCI-TAT-C:

gccaccaggcaatcctgcattgtacataaaaaatctgtatgcagtcagaatttaccatggcacaataaccatttagcagacttcatc
 agataaaaaactcaataactaacgttatctgataaacgttagcattatggaaaccaatctctttatggaaatggaaagggttagtagctta
 cttacataaaaaactgattgtccccaccgataaagaagcatctaaagcatgggacgctatccaccccgitttcattttggcttac
 aataaaaaccgattgtatcttactatcgatttcggagaaaaactcattcaaccagtgaggctcaggcaggcttaaagtaaatt
 agatttcactggctggcgtactgtggagtccttaataacgatcttgcggaaaatcgagagatgaccataatgcaaccaatccctcg
 atggactcaagacagcattggcgtttagtgcataagtcgatagtattcgtttaaagcgcctctaattgtgagtcagggtgaaatc
 tatatcgaccgtattatgtttctgtcgtatgtatcgatgtatcgatccatcaatgttttttttttttttttttttttttttttttt
 tcacaacgtaaagccacaactacctgtacacccgtggaaaatttagccgttgcattatccgcacacttataatgaaatgtcgagg
 gtggaaaagagacaaacctcgattaaagagaatatcagcaattaaatggacttgcatttttttttttttttttttttttttttttt
 gtggaaacgcaggcagacatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtat
 aattatgttatttttagtaattacacgacattaatgttataattagccgttgcgtggaaaagatcccacacaaaaggcgcaacta
 aagcagatgtacttataatgacaaagcatttttagatcaaggcttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
 gttctctgtgggtatattccacgttattaaatgtctgtatgcactaaagaagcgaacctacaaactcaagtttttttttttttt
 tcacgtgagttaaaatgttt
 attactactagagcctgtatgtatcaacttagttataactttcagccattatatcactggcgttgcgttgcgttgcgttgcgt
 gtggtaaagatgtttacgcctgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtat
 cttagcttatt
 acagtaatccagaagttggattaccgctgcaggaaacaccctttaactcaccttcgtttaaatcagtcgttgcgttgcgttgcgt
 tgccatgtctgaaaatcatgcctgtataaaacacttgcattttatctgtcgttgcgttgcgttgcgttgcgttgcgttgcgt
 ggagaaaactattacaccagcgttcttactcaagggttctatgcctttaatggcgttgcgttgcgttgcgttgcgttgcgt
 acactgaaagttataacaccaatgttggcatctgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtat
 aatagtggatgtatggctcgagcttcacaggctatcagcaagaagggtggattggatagaatgccagggcaaccactatccac
 ctccctcttaaagacttagacagtcttaacccatcttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
 aatatggcatgtatggcatttcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
 ataatcacttaatt
 attaaatacccttggattaatggacaaaagatagaaaacatgccttataactcaacacttcacaagggttgcgttgcgttgcgt
 gcaatggttacttaattactcaagcagaaaaatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtat
 gaaggaaaacttagctcggcatggatcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
 gaaaaatggagatggcacaatggatggatggatggatggatggatggatggatggatggatggatggatggatggatggat
 ataaactcagcaatgtacggatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgt
 gatgactcatcgtacaaaagacacttt
 atcaatgtcacttacatggcaatgtcaatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgtatgt
 ttacgagttacttggattccacaagaaatcaaactctgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
 ggtcgtaaaaagcgtcgtcaacgtcgtcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt

SEQ ID NO. 46: V. Amino acid sequence for ABCI-TAT-C:

atsnpafdpknlmqseiyhfaqnnpladffssdknsiltlsdkrsimgnqsllwkwgssflhkklivptdkeaskawgrsstpv
 fsfwlynekpidgyltidfgeklistseaqagfkvldftgwrvgvslndlenremlnatntssdgtqdsigrslgakvdsirfka
 psnvsqgeiyidrimfsvddaryqwsdyqvkvtrsepeiqfhnvkpqlpvtpenlaaidlirlqrlinefvgeketnlaleenisklk
 sdfdalnhtlanggtqgrhlitdkqiiyqopenlnsqdkqlfdnyvilgnytllmfnisrayvlekdptqkaqlkqmyllmtkhlld
 qgfvkgsalvthhwgyssrwwyistllmsdalkeanlqtqvysllwysrefkssfdmkvsadssldyfntlsrqhlalllep
 dqkrinlvntfshyitgaltqvppggkdglrpdtgatrhegnypgysfpafknasqliyllrdtpfvsgesgwnsllkamvsawi
 ysnpevgplagrhplnspslksvaqgyylamsaksspdktlasiyaisdktqnestaifgetitpaslpqgyafnggafgih
 wqdkmvtkayntnvwsseiynkdnrirygryqshgvaqivsngsqlsqyqqegwdwnrmpgattihplkdldspkphilm
 qrgergfsgtsslegqygmmafdliyanlerfdpnftakksvlaadnhlifigsninssdknkvetlfqhaitptlntlwingqki

enmpyqtlqqgdwlidsngngylitaekvnvsrqhvsokenrqrptegnfssawidhstrpkdasyeymvldatpekm
gemaqkfrennglyqvrlkdkdvhieldklsnvtgyafyqpasiedkwikkvnkpaivmthrqkdtlivesavtpdlnmtrqkaa
tpvtinvtingkwqsadknsevkyqvsgnteltftsyfgipqekiqlsplgrkkrrqrrppqc

SEQ ID NO. 47 Nucleotide sequence for chondroitinase ABCI-nΔ20

gc acaaaataac

61 ccattagcag acttctcatc agataaaaac tcaataactaa cgttatctga taaacgtac
121 attatggaa accaatctt ttatggaaa tggaaagggtg gtagtagctt tactttacat
181 aaaaaactga ttgtccccac cgataaagaa gcatctaaag catggggacg ctcatccacc
241 cccgtttct catttgcgtt ttacaatgaa aaaccgattt atggttatct tactatcgat
301 ttccggagaaa aactcattt aaccagttag gtcaggcag gctttaaagt aaaatttagat
361 ttcaactggct ggctactgtt gggagtcgtt ttaataacg atctgaaaa tcgagagatg
421 accttaaatg caaccaatac ctccctgtat ggtactcaag acagcattgg gcgttcttta
481 ggtgctaaag tcgatagttt tcgtttaaa ggcgcctcta atgtgagtca gggtaatc
541 tatatcgacc gtattatgtt ttctgtcgat gatgtcgctt accaatggc tgattatcaa
601 gtaaaaaactc gcttattcaga acctgaaaatt caatttccaca acgttaagcc acaactaccc
661 gtaacacccg aaaaatttagc ggccattgtat ttatttcgc aacgtctaat taatgaattt
721 gtcggaggtg aaaaagagac aaacctcgca tttagaagaga atatcagcaa attaaaaaagt
781 gatttcgtat ctcttaatac tcacacttta gcaaatggtg gaacgcaagg cagacatctg
841 atcaactgata aacaaatcat tattttatcaa ccagagaatc ttaactctca agataaaacaa
901 ctatttgata attatgttat tttaggtat tacacgacat taatgtttaa tattagccgt
961 gcttattgtgc tgaaaaaaga tcccacacaa aaggcgcac aaaaagcagat gtactttata
1021 atgacaaagc atttatttaga tcaaggctt gttaaaggaa gtcgtttagt gacnacccat
1081 cactggggat acagttctcg ttgggtgtt attccacgt tattatgtc tgatgcacta
1141 aaagaagcga accttacaaac tcaagtttat gaticattac tggtgttac acgtgagttt
1201 aaaaagtagtt ttgatatgaa agtaagtgt gatagtctg atctagatta ttcaataacc
1261 ttatctcgcc aacatttagc ttattacta cttagagctg atgtcaaaa gcgtatcaac
1321 ttagttataa ctttcagcca ttatatcact ggccgtttaa cgcaagtgc accgggttgt
1381 aaagatgggtt tacggccctga ttggtagc tggcgcacatg aaggcaacta tccgggtacta
1441 tctttccag cttttaaaaa tgcctctcg ttatttttatt tattacgcgta tacaccattt
1501 tcagtggtt aaagtgggtt gaatagccgtt aaaaagcga tggtttcagc tggttactac
1561 agtaatccag aagtggattt accgcgttca ggaagacacc ctcttaactc accttcgttta
1621 aaatcagtcg ctcaaggctt ttactggctt gccatgtctg caaaatcatc gcctgataaaa
1681 acacttgcattt ctatttactt tggtgtttagt gataaaacac aaaaatgaatc aactgttatt
1741 ttggagaaa ctattacacc agcgttttta cctcaagggtt tctatgcctt taatggcggt
1801 gctttggta ttcatcggtt gcaagataaa atggtgacac tgaaagctta taacaccaat
1861 gtttggcat ctgaaattta taacaaagat aaccgtttagt gccgttacca aagtcatgg
1921 gtcgtcaaa tagtggat tggctcgacg ctttcacagg gctatcagca agaagggttgg
1981 gattggaaata gaatgccagg ggcaaccactt atccacccctc ctcttaaaga ctttagacagt
2041 ccttaaacccctt ataccttaat gcaacgttga gagcgtggat tttagcggaaac atcatccctt
2101 gaaggtcaat atggcatgtt ggcattcgat ttattttatc ccgcctaatct tgagcggttt
2161 gatccttaattt tcacttgcgaa aaagagtgtt tttagccgtt ataatcactt aatttttatt
2221 ggttagcaataa taaatgttagt tgataaaaaat aaaaatgtt aacgcacccatttccacat
2281 gccattactc caacattaaa tacccttgg attaatggac aaaagataga aacatgcct
2341 tatcaaaacaa cacttcaaca aggtgttgg ttaattgtata gcaatggcaa tggttactta
2401 attactcaag cagaaaaaagt aaatgtttagt cgccaaacatc aggttccgc gggaaaataaaa
2461 aatcgccaaac cgacagaagg aaactttagc tcggcatggta tcgatcacag cactcgcccc
2521 aaagatgcca gttatggat ttaggtttttttagatgca cacctgaaaa aatggggagag
2581 atggcacaataa aattccgttga aaataatggg ttatatcagg ttcttcgttaa ggataaaagac

2641 gttcatatta ttctcgataa actcagcaat gtaacgggat atgccttta tcagccagca
2701 tcaattgaag acaaatggat caaaaagggtt aataaacctg caattgttat gactcatcgaa
2761 caaaaagaca ctcttattgt cagtgcagtt acacccgtttaaaatgac tcgccccaaaa
2821 gcagcaactc ctgtcaccat caatgtcacg attaatggca aatggcaatc tgctgataaaa
2881 aatagtgaag tgaaaatatca gggttcttgtt gataacactg aactgacgtt tacgagttac
2941 ttggatttc cacaagaat caaaactctg ccactccctt ga

SEQ ID NO. 48: II. Nucleotide sequence for ABCI-pΔ60:

SEQ ID NO. 49: Nucleotide sequence for TAT

ggtcgtaaaaagcgtcgtcaacgtcgctgcctcataatgc

(SEQ ID NO. 50) Amino acid sequence for a TAT peptide

grkkrrqrrrppqc