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Abstract

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Brief Communications

An Inhibitor of Serine Proteases, Neuroserpin, Acts as a Neuroprotective Agent in a Mouse Model of Neurodegenerative Disease

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Various studies suggest that proteolytic activity may be involved in a number of neurodegenerative disorders, including stroke and seizure. In this report, we examined the role of tryptic serine proteases, plasminogen activators (PAs), in the evolution of a neurodegenerative disease. Transgenic mice overexpressing an axonally secreted inhibitor of serine proteases (neuroserpin) were crossed with mice characterized by a “dying-back” motor neuron disease [progressive motor neuronopathy (pmn/pmnn)]. Compared with pmn/pmnn mice that showed an increase in PA activity, double mutant mice had decreased PA activity in sciatic nerves and spinal cord; their lifespan was increased by 50%, their motor behavior was stabilized, and histological analysis revealed increased numbers of myelinated axons and rescue of motoneuron number and size. This is the first report showing that a class of serine proteases (PAs) may be involved in the pathogenesis of a motor neuron disease and more specifically in axonal degeneration. Inhibiting serine proteases could offer a new strategy for delaying these disorders.

Key words: motoneurons; axon; spinal cord; plasminogen activator; neuroserpin; pmn mice

Introduction

Proteolytic activity has been suggested to be involved in a number of neurological disorders, including stroke, seizure, and Wallerian degeneration. Recently, it has been shown that plasminogen activators (PAs) are implicated in axonal degeneration in a mouse model of multiple sclerosis, experimental allergic encephalomyelitis (Lu et al., 2002; East et al., 2005), and other studies have shown that PAs are activated during Wallerian degeneration (Bignami et al., 1982; Siconolfi and Seeds, 2001).

PAs [tissue PA (tPA) and urokinase PA (uPA)] catalyze the conversion of serum plasminogen to plasmin, the active fibrinolytic enzyme, and they are expressed in the CNS in neurons and glia. PAs have been shown to play a role in axonal elongation and nerve regeneration (Siconolfi and Seeds, 2001). It has been proposed that deregulation of the PA system could aggravate neurodegenerative processes in many pathological situations. For example, PAs and in particular tPA mediate neuronal degeneration in models of stroke, epilepsy, and in Alzheimer’s disease (for review, see Melchor and Strickland, 2005).

In this report, we determined whether PAs could be involved in the neurodegenerative process of axons and cell bodies of motoneurons in a mouse model of a neurodegenerative disease called pmn, for progressive motoneuropathy. pmn/pmnn mice are characterized by a “dying-back” motor neuronopathy (Schmalbruch et al., 1991); the pathophysiology begins in the axon terminals by an unknown process, followed by a degeneration of the motoneuron cell bodies. We first show that the activity of PA is increased in sciatic nerves and spinal cord of pmn/pmnn mice compared with controls. We then crossed pmn mice with transgenic mice that overexpress an inhibitor of PAs, i.e., neuroserpin. In these double mutant mice, the lifespan, the motor behavior, the number of surviving motoneurons in the spinal cord and cranial nuclei, and the number of surviving myelinated axons are increased.

Materials and Methods

Animal models

pmn mice were obtained from the laboratory of J. L. Guénet (Institut Pasteur, Paris, France). Mice that overexpressed neuroserpin [TgN (hsNSwt)682 line called ThyNs mice] (Cinelli et al., 2001) were crossed with pmn/+ to generate pmn/+;ThyNs progeny (F1); these mice were crossed to generate the F2 line that contained pmn/pmnn;ThyNs mice. Mice genotypes were screened by PCR. All of the experimental procedures were approved by the Ethical Committee for Animal Experimentation of the Geneva Veterinary Office.

Behavioral analysis

Spontaneous locomotor activity was recorded in a photocell activity chamber (Letica model LE 8811; Biosb, Chaville, France), and data were analyzed using the program SeDaCom32 (Biosb).

Analysis of myelinated axons in the facial and phrenic nerves

Nerves were removed, postfixed, dehydrated, sectioned, and stained with Borsac blue as described previously (Ferri et al., 2003).
Histological analysis of fluorescently labeled sciatic motoneurons
Labeling was performed as described previously (Perrelet et al., 2004). The fluorescent retrograde tracer fluorogold (Fluorochrome, Englewood, NJ) was injected into the gastrocnemius muscle, and animals were killed 1 week later (at 40 d of age). The spinal cords were sectioned at 30 μm on a cryostat. Labeled motoneurons were counted in the ventral horn of the spinal cord in segments L3-L6.

Counting of motoneurons
In spinal cord. Mice were perfused at 12, 20, and 40 d of age. The spinal cords were cryoprotected in 20% sucrose, sectioned at 14 μm on a cryostat, and stained with cresyl violet.
In cranial nuclei. After perfusion the brains were removed, dehydrated in alcohol, embedded in paraffin, sectioned at 8 μm on a microtome, and stained with cresyl violet.
In both cases, only motoneurons with a clearly identifiable nucleus were counted on both sides of the spinal cord in every fifth section; the values were multiplied by five to give the final estimate of the number of motoneurons.

Analysis of the cell body surface of motoneurons
The cross-sectional area of the motoneurons was measured with RoboSoftware (Palm Microlaser Technologies, Bernried, Germany).

Zymographic assay for PAs
Mice were killed and perfused with PBS/liquemine. Sciatic nerves were removed, crushed with a Polytron, and centrifuged to eliminate remaining cell fragments. Zymographic assays were done (with 10 μg of protein) according to Vassalli et al. (1984). The optical density of the bands was measured by scanning the gels and determining the number of pixels per surface area (NIH Imagej version 1.30; Wayne Rasband, National Institutes of Health, Bethesda, MD).

Western blotting
Western blotting was performed as described previously (Perrelet et al., 2004). Primary antibodies included anti-β3-tubulin (mouse, 1:500; Chemicon, Hofheim, Germany) and anti-actin (goat, 1:2000; Santa Cruz Biotechnology, Santa Cruz, CA). Secondary antibodies included anti-mouse and anti-goat IgG, horseradish peroxidase linked. The optical density of the bands was measured as described previously for zymography.

Results
PAs are activated in sciatic nerves and spinal cord of pmn/pmn mice
The initial goal of our work was to determine whether PAs can play a role in the pathophysiology of the pmn dying-back motor neuronopathy. These mice develop muscle weakness and atrophy starting at 2 weeks of age and die at ~6 weeks of age. We first observed that PAs were differentially expressed in sciatic nerves of control and mutant mice (Fig. 1A–C). A quantitative analysis by densitometry at 12, 20, and 40 d of age (Fig. 1B, C) showed no difference in PA activity in pmn/pmn mice and control mice at a presymptomatic stage (12 d of age), whereas it was already increased in pmn/pmn mice at 20 d of age. PA activity increased even further at 40 d of age, corresponding to the end stage of the disease.

Differential expression of PA activity was also observed in the spinal cord; gel zymography showed that both tPA and uPA activity were higher in the spinal cord of 20- and 40-d-old pmn/pmn mice versus control mice (Fig. 1D–F). These results suggest that PAs may be candidate factors for playing a role in axonal degeneration and motoneuron loss in pmn/pmn mice. To determine which cell types overexpress PAs in the spinal cord and sciatic nerves, we performed in situ hybridization for both tPA and uPA mRNAs on sections from control and pmn/pmn mice. In spinal cords, an increase in tPA mRNA was observed in pmn/pmn compared with control mice in various cell types (such as neurons and glia) and not only in motoneurons (data not shown). uPA mRNA was undetectable, probably because of the low expression of this serine protease in the spinal cord (Fig. 1D). We did not detect any labeling of uPA and tPA mRNA in sciatic nerves, suggesting that the primary site of PA synthesis was in the spinal cord (data not shown). To study whether immune cells may be involved in axonal degeneration or PA expression, we compared extracts of sciatic nerves and spinal cord from pmn/pmn and control mice by

Figure 1.  A, Gel zymography of PA activity in sciatic nerves of control and pmn/pmn mice at 40 d of age. B, C, Quantification of tPA (B) and uPA (C) activities at 12, 20, and 40 d of age. D, Gel zymography of PA activity in spinal cord of control and pmn/pmn mice at 40 d of age. E, F, Quantification of tPA (E) and uPA (F) activities; values were normalized to the level of controls that were taken as 100% (n = 3; ANOVA analysis, *p < 0.01, **p < 0.001). std, Standard. PA activity in sciatic nerves and in spinal cord was similar between pmn/pmn mice and control mice at a presymptomatic stage (12 d), whereas there was an increase in both uPA and tPA in pmn/pmn mice at a symptomatic age (20 and 40 d).
Neuroserpin overexpression partially rescued the phenotype of pmn/pmnmice

To determine whether inhibiting PAs may modify the neurodegenerative process in the pmn phenotype, we crossed pmn/pmnmice with ThyNs transgenic mice that overexpress an inhibitor of PAs, neuroserpin, under the control of a neuronal promoter (Thyl2) (Cinelli et al., 2001). It has been shown previously that neuroserpin can inhibit both tPA and uPA in vivo (Yepes et al., 2000) and that it is neurally expressed and axonally secreted (Osterwalder et al., 1998). Because PAs are also secreted proteins that function in the extracellular matrix, neuroserpin provides a tool for interfering with PA activity under pathological conditions. Because the neuroserpin and pmn mutant mice were maintained in different strains (C57BL/6 and SV129, respectively), we also analyzed offspring of (pmn/+ × C57BL/6) × (pmn/+ × C57BL/6) matings and showed that disease progression in pmn/pmnmice was not significantly affected by the genetic background (data not shown). No gene dosage effect was observed when comparing pmn/pmnmice carrying one or two copies of the neuroserpin transgene. By zymographic analysis, we confirmed that exogenous neuroserpin overexpression reduced PA activity in sciatic nerve extracts of pmn/pmnmice following the ThyNs transgene was prolonged to ~60 d of age (61.3 ± 5.9; n = 5), i.e., an increase of ~50% (p < 0.001) (Fig. 2B). We did not observe a difference in the lifespan of control and ThyNsmice.

Neuroserpin overexpression did not provide a protection against the massive weight loss seen in pmn/pmnmice. The weight of the pmn/pmnmice stabilized to a value only slightly higher than homozygote pmn/pmnmice before death (Fig. 2C). A photocell activity chamber was used to monitor spontaneous activity (Fig. 2D). In both pmn/pmnmice, and pmn/pmnm;ThyNs, the activity level was dramatically lower than in controls. However, in pmn/pmnm;ThyNs mice, spontaneous activity decreased more slowly and gradually compared with pmn/pmnmice.

In previous studies, we have shown that certain factors such as ciliary neurotrophic factor act to protect both the axon and the cell body and thus increase the lifespan of pmn mice (Sagot et al., 1995a). In contrast, other molecules such as bcl-2 and glial cell line-derived neurotrophic factor were only capable of rescuing the motoneuron cell bodies without preserving the axons; under these conditions, the lifespan of the pmn mice was unaltered (Sagot et al., 1995b, 1996). In the present experiments, we hypothesized that an increase in PA levels may play a role in motoneuron degeneration, and thus inhibition of PAs by neuroserpin should protect motor axons, motoneuron cell bodies, or both. Myelinated axons were counted in the facial motor nerve of 12-, 20-, and 40-d-old pmn/pmnmice, pmn/pmnm;ThyNs, and control littersmates. At a presymptomatic age, there was no difference in the number of myelinated axons between the different groups. As expected, the number of facial axons in pmn/pmnmice was strongly reduced at symptomatic ages (Fig. 3A, B, G). In contrast, axon loss was reduced in pmn/pmnm;ThyNs compared with pmn/pmnmice (Fig. 3B, C, G).

Myelinated fibers in the phrenic nerve that innervate the diaphragm have been reported to be severely affected in pmn/pmnmice, suggesting that these mice die by respiratory failure (Schmalbruch et al., 1991). To determine whether neuroserpin overexpression prevented this loss, myelinated axons were counted in the phrenic motor nerve at the end stage of disease (40 d of age) in pmn/pmnmice, pmn/pmnm;ThyNs, and control littersmates. As described previously (Sagot et al., 1995b; Ferri et al., 2003), the number of phrenic axons was decreased by 45% in pmn/pmnmice; this loss was partially prevented in neuroserpin overexpressing pmn/pmnmice (Fig. 3I).

The pmn disease results from a mutation in the tubulin chaperone E (lbece) gene that leads to a decreased number of microtubules in the axons (Martin et al., 2002). To determine whether neuroserpin overexpression could stabilize microtubules, we examined the level of β3-tubulin as a quantitative index of microtubule abundance. In the sciatic nerve, at 40 d of age and as determined by Western blotting, the level of β3-tubulin was decreased by 76% in pmn/pmnmice compared with controls (Fig. 3H), whereas in the presence of neuroserpin overexpression, it was decreased by only 55%. Thus, neuroserpin may contribute to stabilizing or restoring the polymerization of microtubules.

Because pmn/pmnmice are characterized by a loss of motoneurons, we examined whether overexpression of neuroserpin could prevent neuronal degeneration. Motoneurons in the lumbar spinal cord and in four cranial nuclei (oculo/trochlear, trigeminal, facial, and hypoglossal) were counted. Gastrocnemius motoneurons in the spinal cord were selectively labeled by retrograde transport after injection of fluorogold into the gastrocne-
mice (Fig. 3D–F), and the total number of lumbar spinal cord motoneurons were determined after cresyl violet staining (Fig. 3K). Whereas spinal motoneurons degenerated in pmn/pmn mice, neuroserpin overexpression partially prevented the decrease in neuron number (~50% reduction in neuronal loss compared with pmn/pmn mice at 40 d of age) (Fig. 3E,F,J,K). Similarly, whereas four populations of motoneurons in cranial nuclei were decreased in pmn/pmn mice, neuroserpin overexpression partially prevented the decrease in the number of both oculo/trochlear and facial nuclei motoneurons (Fig. 4A) and in the size of facial motoneurons (Fig. 4B–E).

**Discussion**

In this study, we examined the role of PAs, a class of serine proteases, in axonal degeneration and motoneuron loss in the pmn mouse. The mutation in pmn mouse has been localized recently to a tubulin chaperone E gene (tbce) (Bommel et al., 2002; Martin et al., 2002). A mutation of human tbce is responsible for hypoparathyroidism–retardation–dysmorphism syndrome that comprises neurological symptoms (Parvari et al., 2002). Interestingly, a recent study shows that human giant axonal neuropathy, an autosomal recessive neurodegenerative disorder, is related to the degradation of tubulin folding cofactor B (Wang et al., 2005). Thus, pmn mice may provide a useful model for studying diseases related to tubulin modification and not only to motoneuron disease.

The goal of our work is to determine whether PAs can play a role in the pathology of the pmn mouse model. We first observe an increase in PA activity at a symptomatic stage (20 and 40 d of age) in sciatic nerves and spinal cord of pmn/pmn mice compared with control mice. No difference in PA activity is observed at a pre-symptomatic stage, suggesting that PA activation may be acting primarily at a later symptomatic stage, suggesting that PA activation may be acting primarily at a later stage of the disease.

To understand whether this activation of PAs is relevant to mechanisms implicated in axonal degeneration and motoneuron loss, we determined whether the overexpression of an inhibitor of PAs, neuroserpin, could affect the disease progression and symptoms in pmn/pmn mice. Therefore, we crossed pmn/pmn mice with neuroserpin overexpressing mice (ThyNS under the control of a neuronal promoter Thy1.2) and observed an inhibition of PA activity in sciatic nerves of pmn/pmn; ThyNS mice by zymographic analysis, demonstrating that neuroserpin overex-

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**Figure 3.** Neuroserpin overexpression partially protects motor axon and motoneuron loss in pmn/pmn mice. *A–C*, Cross sections of facial nerves in control (A), pmn/pmn (B), and pmn/pmn; ThyNS (C) mice at 40 d of age. Insets, Threefold magnification. In B, there are fewer myelinated facial nerves, and they are less compact than in A or C. *D–F*, Transverse sections of lumbar spinal cords from control (D), pmn/pmn (E), and pmn/pmn; ThyNS (F) mice after fluorogold labeling of motoneurons. *G*, Counts of myelinated axons in facial nerves at 12, 20, and 40 d of age. In pmn/pmn; ThyNS mice (n = 5), the number of myelinated axons was significantly increased (ANOVA analysis, *p* < 0.01) compared with pmn/pmn (n = 4). *H*, Western blot for β3-tubulin in sciatic nerves of control, pmn/pmn, and pmn/pmn; ThyNS mice at 40 d of age and quantification (n = 3). Values were normalized to β-actin; controls were taken as 100% (ANOVA analysis, *p* < 0.01). *I*, Counts of myelinated axons in phrenic nerves at 40 d of age. In pmn/pmn; ThyNS mice (n = 3), the number of myelinated axons was significantly increased (ANOVA analysis, *p* < 0.01) compared with pmn/pmn mice (n = 3). *J*, Counts of fluorogold-labeled motoneurons at 40 d of age. In pmn/pmn; ThyNS mice (n = 4), the number of fluorogold-labeled motoneurons was significantly increased (ANOVA analysis, *p* < 0.01) compared with pmn/pmn mice (n = 4). *K*, Counts of cresyl violet-labeled motoneurons in the lumbar spinal cord at 12, 20, and 40 d of age (n = 3) (ANOVA analysis, *p* < 0.01, **p** < 0.001). Scale bars: *A–C*, 50 μm; *D–F*, 100 μm.
pression inhibits PA activity in axons. The apparition of symptoms in both pmn/pmn mice and in pmn/pmnpmn;ThyNs mice occurs at 2 weeks of age, but we observed a 50% increase in the lifespan of pmn/pmnpmn;ThyNs compared with pmn/pmnpmn mice. Furthermore, the body weight and the motor activity are stabilized at the end stage of disease. These results suggest that PAs play a deleterious role at later stages of the disease and that inhibition by neuroserpin can delay the disease progression and increase the lifespan of pmn/pmnpmn mice.

Moreover, we show that neuroserpin overexpression partially protects myelinated motor axons of facial and phrenic nerves and protects cranial motoneurons in the oculo/trochlear and facial nuclei as well as motoneurons in the spinal cord of pmn/pmnpmn mice. Cranial motoneurons also exhibit a selective vulnerability in different mouse models of amyotrophic lateral sclerosis and in human patients; a selective protection of cranial nuclei has been observed previously in pmn/pmnpmn;Wlds mice (Ferri et al., 2003). A number of hypotheses have been put forward to explain why motoneurons do not display a uniform degeneration, including a differential expression of glutamate receptors and glutamate transporter subtypes. Motoneurons in different cranial nuclei may also exhibit a differential sensitivity to PAs and/or neuroserpin or a differential level of expression of these genes.

Several mechanisms have been proposed to explain the role of PAs in axonal degeneration. PAs may promote demyelination because plasmin can directly degrade myelin basic proteins (Cammer et al., 1978). Furthermore, plasmin is the key initiator of the matrix metalloproteinase (MMP) activation cascade and has been documented to play an important role in the breakdown of myelin membranes (for review, see Cuzner and Opdenakker, 1999). Moreover MMP7, MMP9, and MMP12 are activated in optic nerve axons during Wallerian degeneration (Hughes et al., 2002). PAs could also play a direct role in apoptosis of motoneurons. Indeed, it has been shown that tPA plays a role in neuronal apoptosis (Flavin et al., 2000) and that neuroserpin protects neurons from ischemia-induced death (for review, see Galliciotti and Sonderereg, 2006). One prevailing hypothesis is that proteolytic disruption of the basement membrane by tPA may be the trigger that initiates apoptosis. Thus, the capacity of neuroserpin to block tPA-induced degradation of the basement membrane may explain the ability of neuroserpin to reduce neuronal death.

Interestingly, in our mouse model, we observed that neuroserpin overexpression partially rescues the microtubule loss in sciatic nerves of pmn/pmnpmn mice. It has been shown previously that tPA can play a role in microtubule destabilization in hippocampal neurons (Medina et al., 2005). Together, these results suggest that tPA could play a role in the microtubule loss observed in pmn/pmnpmn mice and that neuroserpin overexpression may protect pmn/pmnpmn mice by controlling tPA and its effect on microtubule abundance. Several studies have shown that tPA can play a protective role in axonal regeneration and functional recovery after sciatic nerve injury (Siconolfi and Seeds, 2001; Zou et al., 2006). These results are not incompatible with our own observations because, in experimental paradigms using nerve crush, it is evident that extracellular debris must be removed by some type of proteolytic system before axonal regeneration can occur. Our results favor the implication of PAs in axonal degeneration in a dying-back motor neuronopathy and possibly in the pathogenesis of motoneuron degeneration. Similarly, it has been shown that PAs are implicated in axonal degeneration in a mouse model of multiple sclerosis, experimental allergic encephalomyelitis (Lu et al., 2002; East et al., 2005). Thus, PAs could be implicated in axonal degeneration under different pathological conditions.

It has been shown previously that uPA is activated in wobbler mouse muscle, another mouse model of motoneuron disease (Blondet et al., 1992), but our results are the first to implicate PAs in axonal degeneration and motoneuron loss in a dying-back motor neuronopathy. More importantly, they also show that increased expression of an inhibitor of these enzymes affects the outcome of the disease. It will be interesting to determine whether the attenuation of PA activity may offer a new strategy for delaying axonal degeneration and motoneuron loss in neurodegenerative disorders.

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**Figure 4. A** Number of motoneurons in cranial nuclei in control, pmn/pmnpmn, and pmn/pmnpmn;ThyNs at 40 d of age (± SEM). The number of animals examined is in parentheses. *p < 0.05 by ANOVA analysis between pmn/pmnpmn and pmn/pmnpmn;ThyNs mice. B–D, Micrographs of the facial nucleus in control (B), pmn/pmnpmn (C), and pmn/pmnpmn;ThyNs (D) mice. Scale bar, 50 µm. **E,** Size of facial motoneuron cell bodies at 40 d of age (Control, pmn/pmnpmn, pmn/pmnpmn;ThyNs). *p < 0.01, ANOVA analysis.


