The plasminogen activator system modulates sympathetic nerve function

Ulrich Schaefer,1 Takuji Machida,1 Sandra Vorlova,2 Sidney Strickland,2 and Roberto Levi1

1Department of Pharmacology, Weill Medical College of Cornell University, New York, NY 10021
2Laboratory of Neurobiology and Genetics, The Rockefeller University, New York, NY 10021

Sympathetic neurons synthesize and release tissue plasminogen activator (t-PA). We investigated whether t-PA modulates sympathetic activity. t-PA inhibition markedly reduced contraction of the guinea pig vas deferens to electrical field stimulation (EFS) and norepinephrine (NE) exocytosis from cardiac synaptosomes. Recombinant t-PA (rt-PA) induced exocytotic and carrier-mediated NE release from cardiac synaptosomes and cultured neuroblastoma cells; this was a plasmin-independent effect but was potentiated by a fibrinogen cleavage product. Notably, hearts from t-PA–null mice released much less NE upon EFS than their wild-type (WT) controls (i.e., a 76.5% decrease; P < 0.01), whereas hearts from plasminogen activator inhibitor-1 (PAI-1)–null mice released much more NE (i.e., a 275% increase; P < 0.05). Furthermore, vasa deferentia from t-PA–null mice were hypersensitive to EFS (P < 0.0001) but were normalized by the addition of rt-PA. In contrast, vasa from PAI-1–null mice were much more responsive (P < 0.05). Coronary NE overflow from hearts subjected to ischemia/reperfusion was much smaller in t-PA–null than in WT control mice (P < 0.01). Furthermore, reperfusion arrhythmias were significantly reduced (P < 0.05) in t-PA–null hearts. Thus, t-PA enhances NE release from sympathetic nerves and contributes to cardiac arrhythmias in ischemia/reperfusion. Because the risk of arrhythmias and sudden cardiac death is increased in hyperadrenergic conditions, targeting the NE-releasing effect of t-PA may have valuable therapeutic potential.
enrichment; references 19, 20). Stimulation of PC-12 cells and primary bovine adrenal chromaffin cells with nicotine, KCl, and BaCl₂ results in a prominent corelease of catecholamines and t-PA (19, 20). Accordingly, sympathoadrenal activation may be an important physiologic mechanism for the rapid release of t-PA.

Other than fibrinolysis, the functional significance of the release of t-PA is presently not fully understood, but an involvement in extracellular matrix degradation, angiogenesis, vascular remodeling, and even plaque rupture has been proposed (22–24). In the brain, this serine protease has been shown to play diverse roles in addition to its thrombolytic activity and participate in long-term potentiation, excitotoxicity, and postischemic neurodegeneration (25).

The aim of this study was to investigate whether t-PA is involved in sympathetic neuronal function and transmitter release. We report that t-PA participates in peripheral sympathetic transmission by promoting norepinephrine (NE) release from sympathetic terminals and, in doing so, is likely to contribute to cardiac arrhythmias initiated by NE in ischemia/reperfusion.

**RESULTS**

t-PA inhibition attenuates the contractile response of guinea pig vas deferens to electrical field stimulation

The isolated vas deferens, with its very dense sympathetic innervation, has been used extensively as a model to investigate drug effects at pre- and postsynaptic sites of the sympathetic junction (26, 27). As a first assessment of a possible role of t-PA in sympathetic transmission, we investigated whether the antagonism of t-PA with its synthetic inhibitor 2,7-Bis(4-amidinobenzylidene)cycloheptan-1-one dihydrochloride, t-PAstop (28), would affect the response of the vas deferens to sympathetic nerve stimulation. Sympathetic nerves were stimulated by electrical field stimulation (EFS; 0–64 Hz, supramaximal voltage; stimulation duration of 15 s at 5-min intervals, with pulses lasting 1 ms each). Typically, the contractile response of the vas deferens to EFS consists of two phases: an initial spike (A; purinergic) followed by a plateau (B; adrenergic). Peak response amplitudes (means ± SE [error bars]; n = 4) are expressed as percentages of the response to 40 mM K⁺. Responses were recorded either in the absence or presence of 1 and 10 μM tPAstop. (C, top) Release of endogenous NE from guinea pig heart synaptosomes by depolarization with 3–100 mM K⁺. Points are mean increases in NE release above basal level (± SE; n = 10; EC₅₀ = 32.8 mM). (bottom) Concentration response curves for the inhibition of NE exocytosis elicited by depolarization with 100 mM K⁺ by t-PAstop or rPAI-1. Equieffective inhibitory concentrations of t-PAstop and rPAI-1 were predetermined in a photometric assay of t-PA activity inhibition (IC₅₀ 1.2 μM tPAstop and 9.2 nM rPAI-1; not depicted). Points (means ± SE; n = 4) are expressed as the percent inhibition of NE release by 100 mM K⁺.
Neuroblastoma cells ([3H]NE). Points are means (± SE; n = 4–8) of increases in NE release above baseline. *, P < 0.05 versus baseline; **, P < 0.01 versus baseline.

Concentration response curves for the NE-releasing effects of rt-PA in guinea pig heart synaptosomes (endogenous NE) and human SH-SY5Y neuroblastoma cells. Bars are means (± SE; n = 3–10) of percent increases in NE release above baseline. *, P < 0.05 versus rt-PA alone.

Figure 3. Mechanisms of rt-PA–induced NE release in guinea pig heart synaptosomes. (A) Concentration response curves for the NE-releasing effect of rt-PA and plasmin (each at 0.1–10 μg/ml) in guinea pig (GP) heart synaptosomes. As opposed to rt-PA, plasmin did not elicit NE release. Points are means (± SE [error bars]; n = 10 for rt-PA and n = 4 for plasmin). (B) Effects of various inhibitors on the NE-releasing effect of 10 μg/ml rt-PA. Preincubation with the inhibitors reduced the rt-PA–induced NE release from guinea pig heart synaptosomes. 100 nM Ω-conotoxin (α-IICTX), 10 μM BAPTA-AM, 300 nM desipramine (DMI), 30 μM [5-(N-ethyl-N-isopropyl)-amiloride (EIPA), and 1 μM carpipride (HDE642) each significantly inhibited rt-PA–induced NE release. In contrast, 0.2 μM α2-antiplasmin (α2-AP) did not affect the NE-releasing effect of rt-PA. Bars are means (± SE; n = 3–12) of percent increases in NE release above baseline. *, P < 0.05 versus rt-PA alone.

The administration of recombinant t-PA elicits NE release in guinea pig heart synaptosomes and neuroblastoma cells

Because the inhibition of t-PA activity was associated with a marked inhibition of sympathetic function and NE exocytosis (Fig. 1), it was possible that an increase in t-PA availability would yield the opposite effect. To address this possibility, we incubated guinea pig heart synaptosomes with 0.1–10 μg/ml of recombinant t-PA (rt-PA). rt-PA elicited a dose–dependent increase in the release of NE (EC50 = 1.17 ± 0.15 μg/ml; Fig. 2 A). In a second set of experiments, we preincubated cardiac synaptosomes with a well-known activator of t-PA, cyanogen bromide–digested fibrinogen (CNBr-F; reference 32). Preincubation with 200 μg/ml CNBr-F greatly enhanced t-PA–induced NE release in cardiac synaptosomes. The amount of NE released by rt-tPA (1 μg/ml) increased ~3.5-fold when synaptosomes were preincubated with CNBr-F (Fig. 2 B).

We subsequently extended this investigation to the neuroblastoma cell line SH-SY5Y, which is an ideal model of the sympathetic neuron (33). rt-PA induced [3H]NE release in a concentration range similar to that effective in cardiac synaptosomes (Fig. 2 C). Furthermore, CNBr-F potentiated the NE-releasing effect of t-PA in SH-SY5Y cells to an extent similar to that observed in cardiac synaptosomes (Fig. 2 D).

Mechanisms of rt-PA–induced NE release in guinea pig heart synaptosomes

Next, we explored the mechanisms of t-PA–induced NE release. Initially, we questioned whether the release of NE by t-PA is a primary effect or secondary to plasmin formation.
For this, we first determined whether plasmin has the capacity of releasing NE. At concentrations of 0.1–10 μg/ml, plasmin failed to elicit NE release from guinea pig heart synaptosomes, whereas in the same concentration range, rt-PA caused an ∼5–22% increase in NE release above basal level (i.e., 72% of the release obtained with 100 mM K⁺; Fig. 3 A). To further test whether the NE-releasing effect of t-PA is independent of the generation of plasmin, we preincubated cardiac synaptosomes with 0.2 μM of the plasmin inhibitor α₂-antiplasmin (8) and exposed them to the highest effective dose of rt-PA (10 μg/ml). α₂-Antiplasmin did not affects the rt-PA–induced NE release (Fig. 3 B).

We next investigated whether the NE-releasing effect of t-PA is a Ca²⁺-dependent exocytotic process. Guinea pig heart synaptosomes were incubated with 100 nM of the N-type Ca²⁺ channel blocker ω-conotoxin (34) and were challenged with 10 μg/ml rt-PA. ω-Conotoxin inhibited rt-PA–induced NE release by ∼30% (Fig. 3 B). Pretreatment of cardiac synaptosomes with 10 μM of the intracellular Ca²⁺ chelator BAPTA-AM (35) also markedly reduced (by ∼50%) the rt-PA–induced NE release. This suggested that t-PA may release NE by an exocytotic mechanism that is dependent both on the influx of extracellular Ca²⁺ and an increase in intracellular Ca²⁺. However, the NE-releasing effect of 10 μg/ml rt-PA was also attenuated (by ∼40%) when cardiac synaptosomes were preincubated with 300 nM of the NE transporter inhibitor desipramine (36) and with each of two Na⁺/H⁺ exchange inhibitors, 5-(N-ethyl-N-isopropyl)amiloride (EIPA; 30 μM; ∼80% inhibition) and cariporide (compound HOE642; 1 μM; ∼40% inhibition; reference 36). These findings suggested that in addition to eliciting NE exocytosis, t-PA is also likely to elicit NE release via the NE transporter operating in an outward direction (37).

t-PA gene deletion attenuates and PAI-1 gene deletion potentiates the contractile response of mouse vas deferens to EFS

Because our findings in guinea pig preparations and the neuroblastoma cell line indicated a role for t-PA in sympathetic nerve activity, we further tested this possibility in mice lacking either the t-PA (t-PA−/−) or PAI-1 gene (PAI-1−/−). The contractile responses of vasa deferentia isolated from control mice (WT) to EFS had both purinergic and adrenergic phases that were characterized by a progressive increase in tension with increasing frequencies of stimulation (4–64 Hz). The contractile responses to EFS were greatly diminished in vasa isolated from t-PA−/− mice (P < 0.0001 vs. WT; purinergic and adrenergic phases); in contrast, there were greater increases in tension in response to EFS in vasa isolated from PAI-1−/− mice (P < 0.05 vs. WT; purinergic phase; Fig. 4, A and B) Frequency response curves for the contractile responses of the isolated mouse vas deferens to electrical field stimulation (EFS; 0–64 Hz, supramaximal voltage; every 1 ms for 15 s). Peak response amplitudes of both purinergic and adrenergic phases (means ± SE [error bars]; n = 6–9) are expressed as percentages of the response to 80 mM K⁺. Vasa deferentia isolated from t-PA−/− mice developed markedly less tension in response to EFS than vasa from WT control mice. In contrast, vasa from PAI-1−/− mice developed more tension than vasa from WT mice. Vasa from plasminogen−/− mice developed the same tension as vasa from WT mice. (C) Coronary NE overflow from isolated mouse hearts in response to EFS (0–9 Hz; 5 V for a duration of 60 s, with pulses of 2 ms each). Hearts were perfused with buffer containing 0.1 μM desipramine, 0.1 μM rauwolscine, 1 μM atropine, and 10 μM hydrocortisone. NE overflow was significantly smaller in hearts from t-PA−/− mice than in hearts from WT mice, whereas it was significantly greater in PAI-1−/− hearts. Points are means ± SE (n = 6–9) of x-fold increases in NE overflow above basal levels. (D) K⁺-induced NE exocytosis in mouse heart synaptosomes. Points are means ± SE (n = 12–16) of increases in NE release above basal levels. Synaptosomes isolated from hearts of t-PA−/− mice released significantly less NE in response to K⁺ than synaptosomes from WT hearts. In contrast, synaptosomes from PAI-1−/− hearts released greater amounts of NE than synaptosomes from WT hearts, whereas NE exocytosis in synaptosomes from plasminogen−/− mice was not different from that of synaptosomes from WT hearts.
A and B). Notably, in vasa deferentia from plasminogen-null mice, the contractile responses to EFS were not different from WT controls (Fig. 4, A and B), supporting the notion that the modulatory function of t-PA at sympathetic junctions is independent of plasminogen availability.

**t-PA gene deletion attenuates and PAI-1 gene deletion potentiates NE exocytosis in the mouse heart**

In isolated Langendorff-perfused mouse hearts, EFS (3–9 Hz) elicited a frequency-dependent increase in NE overflow over baseline (Fig. 4 C). NE overflow was greatly diminished in hearts isolated from t-PA−/− mice as compared with WT control hearts (a 76.5% decrease; P < 0.01). In contrast, NE overflow was markedly increased in hearts isolated from PAI-1−/− mice (a 275% increase over WT; P < 0.05; Fig. 4 C).

Because the experiments with Langendorff-perfused mouse hearts suggested that the absence of t-PA leads to a decrease in NE exocytosis from sympathetic nerves, we extended our observations to the cardiac synaptosome model. Depolarization of mouse heart synaptosomes with increasing K+ concentrations (30, 50, and 100 mM) resulted in a concentration-dependent increase in NE release (Fig. 4 D). NE release was greatly reduced in synaptosomes isolated from t-PA−/− mouse hearts but were markedly enhanced in PAI-1−/− synaptosomes (Fig. 4 D). K+ induced NE release in synaptosomes isolated from plasminogen−/− mice was not different from that of their WT controls (Fig. 4 D).

Similar to NE, K+ induced (100 mM) ATP release (assessed with a luciferin-luciferase assay; reference 38) was 4.4-fold (P < 0.01) greater in synaptosomes from WT mice than in those from t-PA−/− mice. Furthermore, 10 μM t-PA stop markedly reduced (by 45.8%; P < 0.05) the K+ induced ATP release in synaptosomes from WT mice.

**The administration of rt-PA restores the contractile response of vasa deferentia isolated from mice lacking t-PA**

We questioned whether supplying exogenous t-PA to a t-PA-depleted system would restore functional responses to normality. Accordingly, we isolated vasa deferentia from t-PA−/− mice and incubated them with rt-PA. The contractile response of vasa deferentia isolated from WT mice and subjected to EFS (30 Hz) was measured in the absence or presence of 0.6 and 1 μg/ml rt-PA. In the presence of 1 μg/ml rt-PA, both purinergic and adrenergic responses to EFS were significantly increased (P < 0.05), whereas the 0.6-μg/ml concentration was without effect (Fig. 5 A). Notably, when this subthreshold concentration of 0.6 μg/ml rt-PA was added to vasa deferentia isolated from t-PA−/− mice, both purinergic and adrenergic contractile responses were restored to the levels of WT controls (Fig. 5 B).

**The potentiation of sympathetic responses by t-PA is not caused by an action at postjunctival sites**

Because the contractile response of the vasa deferentia to EFS is a composite of pre- and postjunctival events (i.e., neurotransmitter release and postsynaptic effects), we questioned whether t-PA might potentiate sympathetic responses in part by an action at postjunctival sites. Therefore, we assessed whether the postsynaptic effects of sympathetic neurotransmitters (i.e., ATP and NE) would be affected by changes in endogenous t-PA availability. Mouse vasa deferentia were incubated in increasing concentrations of exogenous ATP or NE, and the contractile response was measured. The concentration response curves for the effects of ATP and NE in vasa from t-PA−/− and PAI-1−/− mice were superimposable on the curve obtained from vasa of WT control mice (Fig. 6, A and B). These observations indicated that t-PA modulates sympathetic responses by acting solely at prejunctival sites.

**t-PA promotes NE release and associated arrhythmias in myocardial ischemia/reperfusion**

Langendorff-perfused mouse hearts were subjected ex vivo to 30-min stop-flow global ischemia followed by 30-min reperfusion. Hearts isolated from WT mice released ~90 pg
t-PA potentiates sympathetic responses by an action at prejunctional sites. (A) Noncumulative concentration response curves for the contractile response of mouse vas deferens to the administration of exogenous ATP. Vasa deferentia were isolated from t-PA−/−, PAI-1−/−, and their WT control mice. The maximum contractile response for each increment in ATP concentration occurred within 30 s; ATP was quickly washed out thereafter to prevent receptor desensitization. The following higher ATP concentration was added after 30 min of reequilibration. (B) Cumulative concentration response curves for the contractile response of mouse vas deferens to the administration of exogenous NE. Vasa deferentia were isolated from t-PA−/−, PAI-1−/−, and their WT control mice. The finding that the concentration response curves obtained from vasa deferentia of gene-deleted mice were superimposable on the curves obtained from their WT controls indicates that the lack of t-PA or PAI-1 does not influence postsynaptic responses at sympathetic junctions and that the action of t-PA on sympathetic neurons is limited to presynaptic sites. (A and B) Points are means (± SE; n = 7–10) of maximal contractile responses to ATP (A) and NE (B) expressed as percentages of the response to 80 mM K+. NE/gram of tissue in the first 5 min of reperfusion (Fig. 7 A). NE overflow subsided in the following 5 min. In hearts isolated from PAI-1−/− mice, NE overflow was slightly but not significantly greater than that of control hearts. In contrast, in hearts isolated from t-PA−/− mice, NE overflow during reperfusion was approximately fivefold lower than that of WT control and PAI-1−/− mouse hearts. Preischemia NE overflow was comparable in all hearts. Reperfusion was also associated with the development of ventricular arrhythmias such as ventricular tachycardia (VT), ventricular fibrillation (VF), and premature ventricular contractions (PVCs). The lowest incidence and shortest duration of rhythm disturbances was observed in hearts from t-PA−/− mice (Fig. 7 B). In contrast, PAI-1−/− mouse hearts had the highest number of PVCs. These findings suggested that t-PA may contribute to NE-initiated arrhythmias in ischemia and reperfusion.

**DISCUSSION**

Our findings clearly demonstrate that t-PA plays an important role in peripheral sympathetic responses by promoting neurotransmitter release from sympathetic neurons via an action restricted to presynaptic terminals. As a result, in hyperadrenergic states such as myocardial ischemia/reperfusion, in which excessive NE release is a primary arrhythmogenic factor (39, 40), t-PA participates with NE in arrhythmic cardiac dysfunction.

Although endothelial cells have traditionally been viewed as a major source of t-PA (41, 42), we were intrigued by the report that the stimulation of cardiac sympathetic nerves caused the release of t-PA into the coronary circulation (15) and by the prospect that this t-PA might derive from sympathetic neurons, which can both synthesize it and release it (11, 12). It had also been shown that chemical sympathectomy decreases t-PA release from blood vessels (10) and that t-PA is stored in the same vesicular pool with NE in adrenal chromaffin and PC12 cells, from where it could be released together with NE upon depolarization (19, 20). As all of these studies favored the notion of a dual neuronal and endothelial t-PA release, we questioned what function neuronally released t-PA might have other than a plausible involvement in fibrinolysis, and we hypothesized that t-PA might modulate sympathetic neuronal activity.
We reasoned that if t-PA were to play such a role, it should be possible to uncover it with the use of t-PA inhibitors. For this, we used t-PAstop, a synthetic inhibitor of t-PA (28), and the recombinant form of the high affinity physiological serine protease inhibitor rPAI-1. In the isolated vas deferens of the guinea pig, a prototypical model of sympathetic neuromuscular junction (26, 27), t-PAstop attenuated both the purinergic and adrenergic responses to EFS, demonstrating that a fully functional t-PA is required for a complete sympathetic response and suggesting that neuronal t-PA is likely to potentiate either the release or the effects of the sympathetic neurotransmitters. The first indication that sympathetic nerve terminals are the likely site of this potentiation emerged from finding that each of the two t-PA inhibitors, t-PAstop and rPAI-1, inhibited NE exocytosis as a function of its concentration in guinea pig heart synaptosomes, which is a typical model of sympathetic nerve endings (31).

As these results suggested that t-PA enhances the release of NE elicited by the depolarization of sympathetic nerve terminals, we sought to confirm this view by determining whether the administration of rt-PA to native nerve endings (cardiac synaptosomes) or neuroblastoma cells in culture directly enhanced NE release. We chose to use concentrations of t-PA approximating those clinically achieved for intraarterial thrombolysis (43). We not only found that t-PA indeed promotes NE release but particularly important was discovering that fibrin (CNBr-F), which is known to potentiate the plasmin-generating effect of t-PA (6, 7), also potentiated the NE-releasing effect of rt-PA. Inasmuch as the generation of plasmin is pivotal for the fibrinolytic activity of t-PA, we next assessed whether the NE-releasing effect of rt-PA might also be plasmin dependent. Clearly, it was not. Indeed, the administration of plasmin did not significantly affect basal NE release in guinea pig heart synaptosomes, and the administration of α2-antiplasmin (9) did not modify the NE-releasing effect of rt-PA. Moreover, the contractile response of the vas deferens to EFS and the NE exocytosis from cardiac synaptosomes isolated from plasminogen-deficient mice did not differ from those of WT mice, excluding a role for plasminogen and, thus, for plasmin in these responses. Accordingly, we concluded that t-PA acts directly at sympathetic nerve terminals to release NE by an action that is potentiated by fibrin (CNBr-F) but independent of plasmin formation.

Having determined that sympathetic responses are either attenuated when endogenous t-PA is inhibited or potentiated by the administration of rt-PA, we sought definitive proof of the relevance of t-PA actions in sympathetic neural function in animals lacking t-PA or PAI-1. Consistent with a t-PA-induced promotion of NE release, we found that in tissues isolated from t-PA-null mice, sympathetic responses and NE release were markedly reduced. Conversely, sympathetic responses and NE release were potentiated in tissues from the PAI-1-null mice. The notion that the lack of t-PA was indeed the cause of the depressed sympathetic responses was proven by our reconstitution experiments, demonstrating the restoration of sympathetic responses upon administration of a subthreshold amount of rt-PA in the vas deferens. Most importantly, postjunctional contractile responses elicited by the administration of the neurotransmitters ATP and NE in the vas deferens of mice lacking either t-PA or PAI-1 were indistinguishable from the responses recorded in their WT controls. This clearly excluded the possibility that t-PA might act at a postsynaptic site in its potentiation of sympathetic responses.

NE is known to be released from sympathetic neurons by two modalities: exocytosis and reversal of its transporter in an outward direction (i.e., carrier-mediated release; reference 44). The opening of N-type Ca2+ channels and activation of the Na+/H+ exchanger (NHE) at the axonal membrane are pivotal events initiating these two types of release, respectively (35). NHE activation leads to an increase in intracellular Na+, which is critical for the initiation of carrier-mediated NE release (45). Inasmuch as the inhibition of N-type Ca2+ channels with ω-conotoxin (34) and of NHE with the amiloride derivative EIPA each attenuated the NE-releasing effect of rt-PA in cardiac synaptosomes, both mechanisms of NE release are likely to play a role in the action of t-PA. Given that the intracellular Ca2+ chelator BAPTA-AM and cariporide (compound HOE642, a selective NHE-1 blocker; reference 35) also inhibited the NE-releasing effect of rt-PA, a transient increase in intracellular Ca2+ is probably involved in the activation of NHE-1 and the initiation of carrier-mediated NE release (35). A further indication that the NE transporter is outwardly bound under the influence of t-PA is suggested by the finding that the NE transporter inhibitor desipramine attenuated the NE-releasing effect of t-PA.

How t-PA might modulate Ca2+ entry and intracellular Ca2+ transient and NHE-1 activation, eventually culminating in enhanced NE release, is presently a matter of speculation. The action of t-PA could be initiated by protease–substrate reactions, such as those involved in plasmin formation (23, 43, 46). Indeed, the only known substrate of the remarkably specific t-PA in vivo is a single peptide bond (Arg560–Val561) within the proenzyme plasminogen, which is converted to plasmin (47). However, we found that the neuromodulatory action of t-PA is independent of plasmin formation. Therefore, it is conceivable that either an unknown substrate is activated by t-PA, leading to enhanced transmitter release, or a receptor protein is proteolytically activated by t-PA, thereby disinhibiting/augmenting NE release from nerve endings. In fact, our findings with t-PAstop and CNBr-F suggest that the activity level of t-PA is the major determinant of its neuromodulatory action. So far, a specific t-PA receptor has not been discovered, although different binding proteins for t-PA have been described previously (46, 48, 49). Nevertheless, it is likely that a specific substrate or receptor is implicated because the NE-releasing effect of rt-PA appears to involve mechanisms (i.e., intracellular Ca2+, NHE-1, and NE transporter) known to mediate NE exocytosis and carrier-mediated release, which are all modulated by receptor activation (37).

t-PA has been reported to overflow into the coronary effluent of isolated rat hearts during reperfusion after ischemia.
Also, it was recently shown that myocardial ischemia elicits the release of t-PA in the coronary vasculature of the pig (16, 51). Despite the long-held belief that increased endogenous fibrinolysis is beneficial in cardiovascular disease (52), recent studies in the brain emphasize possible adverse effects of t-PA (25). Notably, t-PA is overexpressed in atherosclerotic coronary arteries (53), and elevated t-PA plasma concentrations correlate with the severity of coronary artery disease (54) and are an independent predictor of myocardial infarction (55, 56). Moreover, arrhythmias are often associated with the thrombolytic use of t-PA in the setting of myocardial infarction (57–61). In hyperadrenergic states such as myocardial ischemia and reperfusion, excessive NE release is a primary arrhythmogenic factor (37, 39, 40), and t-PA is released upon the stimulation of cardiac sympathetic nerves (15). Therefore, it is conceivable that t-PA may participate with NE in the generation of arrhythmias associated with ischemia and reperfusion. In fact, we found that ischemia/reperfusion hearts from t-PA knockout mice released significantly less NE (P < 0.05) and had fewer and briefer instances of VF than WT controls. Thus, endogenous t-PA is likely to contribute to arrhythmias linked to myocardial ischemia and infarction.

The administration of rt-PA for thrombolysis in acute myocardial infarction and stroke is still considered an important treatment option. Despite its beneficial effects on cerebral infarct size, rt-PA is believed to induce neurotoxic effects. In fact, rt-PA infusion has been shown to dramatically increase cerebral infarct size in t-PA−/− mice, whereas infarct size was significantly smaller in untreated t-PA−/− mice than in WT controls (62). Supporting possible adverse cardiac effects of t-PA, we found that hearts from t-PA−/− mice were protected against reperfusion arrhythmias, a phenomenon probably associated with the reduced NE spillover. Moreover, we found that in therapeutically relevant concentrations, rt-PA increased NE release from cardiac synaptosomes and human neuroblastoma cells in culture as a function of its concentration.

In conclusion, we have obtained novel evidence that the endogenous plasminogen activator system plays an important role in promoting sympathetic transmitter release in the heart. This action is independent of plasmin formation. Most importantly, hearts from t-PA−/− null mice released much less NE and had fewer arrhythmias when subjected to ischemia/reperfusion than their WT controls. Thus, the plasminogen activator system, which is characterized by the activity level of t-PA, is involved in the excessive release of NE and associated arrhythmias in myocardial ischemia and reperfusion. Targeting this effect of t-PA may have valuable therapeutic potential not only in myocardial ischemia but also in other hyperadrenergic conditions such as heart failure and hypertension.

**MATERIALS AND METHODS**

**Guinea pig and mouse vas deferens.** 250–400-g male guinea pigs and 4–5-mo-old mice were anesthetized with CO2 and exsanguinated (approved by the Weill Medical College’s Institutional Animal Care and Use Committee).

Vasa deferentia (prostatic portion of guinea pigs and midsegment of mice) were suspended in a 20-ml bath containing Krebs-Henseleit (KH) solution at 37°C aerated with 98% O2 + 5% CO2. Mice deficient in PAI-1, plasminogen, or t-PA (63, 64) were provided by S. Strickland. The background of all mice was C57BL/6. Vasa were equilibrated for 60 min (resting tension of 1 g for guinea pigs and 250 mg for mice). EFS was applied for 15 s every 5 min (pulses of 1 ms at 4–6 Hz; supramaximal voltage). The first contractile phase (PPADS sensitive; purinergic) and second phase (prazosin sensitive; adrenergic) were expressed as percentages of the response to K+ (40 and 80 mM for guinea pigs and mice, respectively). Responses were stable to at least three to four consecutive stimulations. When used, drugs were preincubated for 15 min.

**Guinea pig heart synaptosomes.** Male guinea pigs were killed as indicated above. Hearts were isolated and perfused at constant pressure (40 cm H2O) with oxygenated Ringer’s solution at 37°C for 20 min. Hearts were minced in ice-cold 0.32 M sucrose containing 1 mM EGTA. Synaptosomes (pinched-off sympathetic nerve endings) were isolated as previously described (31) and incubated with various drugs at 37°C in the absence or presence of 0.1–10 μg/ml rt-PA. After centrifugation at 20,000 g for 20 min at 4°C, NE and protein contents were determined in the supernatant and pellet, respectively (31).

**Expression of PAI-1 in Pichia pastoris.** rPAI-1 was prepared by yeast transfection according to the manufacturer (Invitrogen). PAI-1 was purified from culture supernatants by nickel-agarose chromatography; enzyme purity was determined by SDS/PAGE.

**Measurement of PAI-1 activity.** Purified rPAI-1 was diluted in 50 mM Tris-imidazole buffer (300 mM NaCl, pH 8.4) and incubated with 5 μl rt-PA (0.1 mg/ml) for 15 min at room temperature, and the kinetics of substrate cleavage (2 mM pelachrome tPA; Pentapharm) were monitored spectrophotometrically at 405 nm in 96-well microtiter plates for 5 min.

**Cross-reactivity between human reagents and guinea pig proteins.** Experiments were performed to ascertain that recombinant human proteins cross react with guinea pig substrates and, in particular, that rt-PA binds to guinea pig plasminogen and activates it. After purification (i.e., by lysine-Sepharose column separation), guinea pig plasminogen was converted to plasmin by human rt-PA in a concentration-dependent fashion, as demonstrated by SDS-PAGE containing lytic casein bands. This suggested cross-reactivity between human and cavian elements of the plasminogen activator system.

**SH-SY5Y neuroblastoma cell line.** The human neuroblastoma cell line SH-SY5Y, which was provided by T.W. Lovenberg (Johnson and Johnson Pharmaceutical Research and Development, LLC, San Diego, CA), was maintained in a 1:1 ratio of Eagle’s MEM supplemented with 10% FBS, 50 U/ml penicillin, and 50 μg/ml streptomycin at 37°C and 5% CO2. Cells were grown to confluence in six-well plates. [3H]NE release experiments were performed as described previously (65). [3H]NE loading was achieved with Hepes-buffered Na+ Ringer’s solution (50 nM [3H]NE) at 37°C for 60 min. After three washings, 1 and 10 μg/ml rt-PA was added for 10 min at room temperature. When CNBr-F was used, rt-PA was incubated together with CNBr-F at room temperature for 5 min and subsequently added to the samples. Aliquots of the supernatant and cell lysates (after 30 min of 0.3% Triton X-100) were taken from each well and analyzed for [3H]NE content with a scintillation counter.

**Perfusion of mouse hearts ex vivo.** After killing, hearts of gene-inacti- vated mice (tPA−/−, plasminogen−/−, and PAI-1−/−) were excised and cooled in ice-cold KH solution equilibrated with 98% O2 + 5% CO2 (66). An 18-gauge steel cannula was inserted in the aorta, and the heart was perfused at constant pressure (100 cm H2O) with KH at 37°C. Coronary flow was measured by timed collections of the effluent every 5 min. Coronary NE over- flow was measured by HPLC with electrochemical detection as previously
described (66). The detection limit was 180.05 fmol. Only hearts with an initial stable sinus rhythm were considered.

**EFS.** Mouse hearts were perfused with KH containing 1 μM atropine, 10 μM hydrocortisone, 0.1 μM desipramine, and 0.1 μM tauracillic. Two stainless steel paddles were apposed to the heart with their stimulating surface parallel to the interventricular septum. ECG was recorded online. After a 20-min stabilization, three sequential stimulations (3, 6, and 9 Hz at 5 V for a duration of 60 s, with pulses lasting 2 ms) using PowerLab/8SP (ADInstruments) were applied every 15 min. For ischemia/reperfusion, after a 30-min stabilization, mouse hearts were subjected to 30-min stop-flow normothermic global ischemia followed by 30-min reperfusion. ECG was recorded online (1-kHz recording frequency) and analyzed with PowerLab/8SP. The incidence and duration of reperfusion arrhythmias were calculated according to the Lanbeth Conventions (67).

**Preparation of cardiac synaptosomes.** Two hearts per mouse type were perfused for 20 mm as indicated above. Both hearts were minced together in ice-cold 0.32 M sucrose. Synaptosomes were isolated as described above, depolarized with K+ and NE exocytosis was measured as previously described (66).  

**Drugs and chemicals.** Adenosine 5’-triphosphate (disodium salt), NE-HCI, and EIPI were purchased from Sigma-Aldrich. Plasminogen, t-PA, and CNBr-F were purchased from American Diagnostica, Inc., and α2-antiplasmin (human plasma) was purchased from Calbiochem. Re-combinant two-chain tissue t-PA was obtained from Genentech, Inc., and pefachrome t-PA (Pef-a-5037) was purchased from Pentapharm. HOE642 (cariporide) was provided by B.A. Schoelkens (Hoechst Marion Roussel, Frankfurt am Main, Germany). rPAI-1 was prepared by yeast transfection.

**Statistics.** Values refer to means ± SE. One-way analysis of variance followed by a Dunnett’s posttest, one-sample Student’s t test, and unpaired Student’s t test were performed as indicated. P < 0.05 was considered significant.

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