Targeting of a mutant plasminogen activator to circulating red blood cells for prophylactic fibrinolysis.

Sergei Zaitzev, Dirk Spitzer, Juan-Carlos Murciano, Bi-Sen Ding, Samira Tliba, M. Anna Kowalska, Khalil Bdeir, Alice Kuo, Victoria Stepanova, John P. Atkinson, Mortimer Poncz, Douglas B. Cines and Vladimir R. Muzykantov

Program in Targeted Therapeutics, Institute for Translational Medicine and Therapeutics and Department of Pharmacology (SZ, BSD, ST, VRM) and Department of Pathology and Laboratory Medicine (KB, AK, VS, DBC), University of Pennsylvania School of Medicine, Philadelphia, PA 19104, Department of Medicine, Washington University School of Medicine, St Louis, MO 63112, USA (DS, JPA), Centro Nacional de Investigaciones Cardiovasculares, Madrid (Spain) (JCM), Hematology Division, Childrens Hospital of Philadelphia, 191094 (MAK, MP).
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Address for correspondence:

Dr. Vladimir R. Muzykantov, IFEM, University of Pennsylvania School of Medicine, One John Morgan Building, 3620 Hamilton Walk, Philadelphia, PA 19104-6068

Phone: 215-898-9823, FAX: 215-898-0868, e-mail: muzykant@mail.med.upenn.edu

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A list of non-standard abbreviations: tPA – tissue type plasminogen activator; PA – plasminogen activator; PAI-1 – plasminogen activator inhibitor type 1; scFv – single chain antibody variable fragment; GPA – glycophorin A; RBC – red blood cells; CNS – central nervous system; CR1 – human complement receptor type 1; ETI – Erythrina trypsin inhibitor; SFM – serum free cell culture medium; WB – western blot.

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ABSTRACT

Chemical coupling to carrier red blood cells (RBCs) converts tissue type plasminogen activator (tPA) from a problematic therapeutic into a safe agent for thromboprophylaxis. The goal of this study was to develop a more clinically relevant recombinant biotherapeutic by fusing a mutant tPA with a single chain antibody fragment (scFv) with specificity for glycophorin A (GPA) on mouse RBCs. The fusion construct (anti-GPA scFv/PA) bound specifically to mouse but not human RBCs and activated plasminogen; this led to rapid and stable attachment of up to 30,000 copies of anti-GPA scFv/PA per mouse RBC that were thereby endowed with high fibrinolytic activity. Binding of anti-GPA scFv/PA neither caused RBC aggregation, hemolysis, uptake in capillary-rich lungs or in the reticuloendothelial system, nor otherwise altered the circulation of RBC. Over 40% of labeled anti-GPA scFv/PA injected in mice bound to RBC, which markedly prolonged its intravascular circulation and fibrinolytic activity compared to its non-targeted PA counterpart. Anti-GPA scFv/PA, but not its non-targeted PA analog, prevented thrombotic occlusion in FeCl₃ models of vascular injury. These results provide proof-of-principle for the development of a recombinant PA variant that binds to circulating RBC and provides thromboprophylaxis using a clinically relevant approach.
INTRODUCTION

Plasminogen activators (PA, including tissue-type, tPA), proteases generating plasmin, which cleaves fibrin clots and restores perfusion, are used to achieve urgent thrombolysis within a relatively narrow therapeutic time window after thrombosis (Topol et al., 1987; Holvoet et al., 1993). The safety of this approach is limited by the inability of soluble PAs to discriminate newly formed occluding pathological clots from pre-existing mural hemostatic clots, and their efficacy is limited by delay in initiation of treatment, inactivation by plasma inhibitors and inadequate delivery into poorly permeable occlusive clots. Paradoxically, endowing tPA derivatives with higher affinity to clot components (Collen, 1996; Runge et al., 1996) further impairs permeation (Sakharov and Rijken, 1995). Increased dosing and potency also increase the risk of bleeding and collateral damage in the brain.

In theory, prophylactic administration of tPA should benefit patients predisposed to a short-term risk of thrombosis (e.g., immobilized patients post surgery, myocardial infarction or transient ischemic attack). Also, unfavorable pharmacokinetics (circulation time <20 minutes) precludes prophylactic use of tPA. However, coupling tPA to carrier red blood cells (RBCs) fundamentally alters tPA pharmacokinetics, converting it from a problematic therapeutic agent into a safe and effective prophylactic agent (Murciano et al., 2003). Studies in animal models have shown that coupling of tPA to RBCs restricts access of the resultant RBC/tPA both to the CNS and to post-surgical hemostatic clots (Zaitsev et al., 2006; Danielyan et al., 2008). RBC/tPA circulate for many hours, incorporate into and rapidly dissolve newly formed, potentially occlusive clots from within (Murciano et al., 2003). Infusion of RBC/tPA in mice, rats and pigs provide an effective short-term option to prevent thrombotic occlusion in diverse vascular systems, including the cerebral vasculature, without the hemorrhagic and CNS toxicity profile typically seen with free tPA (Murciano et al., 2003; Ganguly et al., 2005; Ganguly et al., 2006; Ganguly et al., 2007; Danielyan et al., 2008; Armstead et al., 2009).
The medical utility of this approach would be enhanced if one could circumvent the need for *ex vivo* conjugation of tPA to the carrier RBCs prior to re-infusion. This goal can be achieved using tPA derivatives endowed with the ability to bind safely to circulating RBCs. Thus, tPA chemically conjugated with a monoclonal antibody specific for human complement receptor type I (CR1, an RBC glycoprotein involved in complement regulation and the clearance of immune complexes) (Fearon et al., 1989), can be safely attached onto circulating RBCs, thereby providing thromboprophylaxis in mouse models of thrombosis (Zaitsev et al., 2006). However, CR1 is a low-abundant glycoprotein with significant variation in expression levels among individuals (500-1,500 copies per human RBC) (Birmingham and Hebert, 2001). Therefore, dosing of anti-CR1/tPA conjugates is limited and may be insufficient in cases of severe thrombosis. Further, there are technical and regulatory hurdles for industrial development and clinical use of drugs chemically conjugated to antibodies.

The goal of this study was to design a more generally applicable approach to produce RBC-targeted fibrinolytics that would also permit coating RBCs with a wider range of drug doses. To achieve this goal, we produced a recombinant tPA derivative fused to a monovalent scFv fragment derived from the monoclonal antibody Ter-119, specific for mouse glycophorin-A (GPA), an abundant and RBC-specific surface molecule (~10^6 copies/RBC) (Kina et al., 2000; Spitzer et al., 2004) similar to its human analogue (Furthmayr and Marchesi, 1976). Previous studies showed that the complement regulatory proteins including decay accelerating factor fused with the Ter-119 scFv enhanced the resistance of RBCs to complement-mediated lysis in vitro (Spitzer et al., 2004) and in vivo (Spitzer et al., 2005). In this study, we fused scFv Ter-119 to a truncated form of mouse tPA containing kringle 2 and the protease domain (truncation of auxiliary tPA domains reduces its clearance and side effects) (Martin et al., 1991; Kohnert et al., 1992). Additional mutations homologous to those in Tenectaplas (K296A, H297A, R298A and R299A) were introduced in the protease domain to confer higher resistance to the plasma inhibitor, PAI-1 (Davydov and Cheng, 2001; Tanswell et al., 2002). Therefore, the PA moiety of
the resultant anti-GPA scFv/PA fusion designed and tested in this study combines the mutations found in the human recombinant tPA mutants Retavase and Tenektaplase (Davydov and Cheng, 2001). In the present study, we tested the *in vitro* activity and pharmacological properties of this anti-GPA scFv/PA fusion protein and demonstrated that this novel agent can be used safely to endow RBCs with high levels of fibrinolytic activity, thus prolonging its bioavailability and providing thromboprophylaxis *in vivo.*
MATERIALS AND METHODS

All chemicals were obtained from Sigma (St. Louis, MO) unless otherwise specified. The following additional reagents were used: fibrinogen from Enzyme Research Labs (South Bend, IN), thrombin from Calbiochem (San Diego, CA), iodogen from Pierce (Rockford, IL), QuickChange Site-Directed Mutagenesis kit from Stratagene (La Jolla, CA), *Drosophila* S2 cells, pMT/Bip/V5-His-A vector and Schneiders S2 cell medium from Invitrogen (Carlsbad, CA), *Drosophila* serum-free medium from Lonza (Walkersville, MD), polymerase chain reaction (PCR) core kit and Rapid DNA ligation kit from Roche (Basel, Switzerland), endonucleases from New England Biolabs (Beverly, MA), Erythrina trypsin inhibitor (ETI) Sepharose from Landing BioTeck Inc.

Proteins were radiolabeled with Na $[^{125}I]$ (Perkin-Elmer, Boston, MA) using the iodogen method according to the manufacturer's recommendations. The free iodine was removed using a Bio-Spin 6 column (Bio-Rad Laboratory, Hercules CA). RBC were obtained from fresh anticoagulated mouse blood and radiolabeled with $[^{51}Cr]Cl_2$ (Perkin-Elmer, Boston, MA), as described (Murciano et al., 2003).

**Cloning of anti-GPA scFv-PA and PA.** We followed the template described by us for fusing plasminogen activators with scFv's using a serine-rich linker peptide (Ding et al., 2005; Ding et al., 2008). Ter-119 is a rat monoclonal antibody (mAb) to mouse glycophorin A (GPA) and has been characterized previously (Kina et al., 2000). The pNscTDdSeY plasmid served as the source of the scFv Ter-119 cDNA sequence and has been described earlier (Spitzer et al., 2004). Briefly, the variable heavy and light chain regions of Ter-119 were joined by PCR with a (GGGGS)$_3$ linker to assemble the scFv Ter-119.

Plasmid (pMT/Bip/V5-His-A expression vector) containing mouse tissue type plasminogen activator (tPA) was prepared as follows. Mouse tPA cDNA (Open Byosystems, Huntsville, AL) and cDNA encoding full-length mouse tPA as well as its truncated form containing only the kringle 2 (K2) and protease (P) domains (Retavase analog) was PCR
amplified using upstream primer 5'-cat ggg agg ttc aga ctc gga gcc cgg tcc tac aga ggc ac -3' for full-length tPA and 5'-cat ggg agg ttc aga ctc cct aag gga aaa agc gag gac-3' for the truncated form to introduce BglII restriction site at the 5' end, and the reverse primer 5'-gag ctg ggc ttc tcg aat cat tgc ttc atg ttg tgc tga atc cag-3' to introduce an XhoI restriction site at 3' end. The PCR products were digested with BglII/XhoI restriction enzymes, purified and ligated into pMT/Bip/V5-His-A vector. Point mutations to convert amino acids 296-299 (KNKR) to AAAA were introduced into both constructs using the QuickChange Site-Directed Mutagenesis Kit (Stratagene) per the manufacturer using the direct primer 5'-cag gct ccc atc ttt gtc gct gcc gca gcg tct cct gga gag aga ttc-3' and the reverse primer 5'-gaa tct ctc tcc agg aga cgc tgc ggc agc gac aaa gat ggg agc ctg-3'.

cDNA encoding anti-GPA scFv was amplified for cloning in the expression vector pMT/Bip/V5-HisA using the upstream primer 5'-cgt acg act agt cag gtg aag ctg cag gag tca gga gga ggc-3', which introduces a restriction site for SpeI at the 5' end, and the downstream primer 5'-ata aga atg cgg cgg cgg cgg aag agc tact ac cgc atg agg aag aag ccc gtt tca gtt cca gct tgg tcc c-3', which appends the sequence of a short peptide linker (SSSSG)₂ and a NotI restriction site at 3' end. The K2 and P domain fragment of mouse tPA was amplified using as primers 5'-ata aga atg cgg cgg cgg cgg aag agc tact ac cgc atg agg aag aag ccc gtt tca gtt cca gct tgg tcc c-3', which introduces a SpeI restriction site at 3' end, and downstream 5'-gag ctg ggc ttc tcg aat cat tgc ttc atg ttg tgc tga atc cag-3' to introduce an XhoI restriction site at 3' end. The anti-GPA scFv/PA was assembled as follows: First, the 2 PCR products were purified and digested with SpeI, NotI, and XhoI, respectively. Second, the 2 digested fragments were ligated and cloned into the SpeI and XhoI sites of the vector pMT/Bip/V5-HisA. Successful cloning was confirmed by restriction analysis of plasmid and by automated sequencing.

Expression and purification of anti-GPA scFv-PA and PA. Drosophila S2 cells were maintained in Schneiders medium (Invitrogen) supplemented with L-Glutamine (Invitrogen), FBS (HyClone) and PenStrep (Invitrogen) and co-transfected using FuGene6 (Roche) with
pMT/Bip/V5-HisA plasmids encoding anti-GPA scFv/PA and PA constructs and pCoBlast (Invitrogen) at a ratio 30:1. Stable transfectants were established by adding Blasticidin (Invitrogen) (25μg/ml). Stable transfectants were then transferred into S2 serum free medium, SFM (Lonza). Protein production was induced by adding CuSO₄ (final concentration 0.5mM). Anti-GPA scFv/PA and PA were purified from cell media by affinity chromatography on ETI sepharose, as described (Heussen et al., 1984). The yield was ~3 and ~5 mg/l medium for anti-GPA scFv/PA and PA, respectively. Proteins were concentrated to a level not exceeding 2mg/ml, separated into aliquots and stored at -80°C until use.

**Biochemical characterization of anti-GPA scFv-PA and PA.** The size and homogeneity of the fusion protein and its plasminogen activator component were analyzed using a 4%-12% SDS-PAGE gradient. For Western blot analysis, the separated proteins were electrotransferred to a nitrocellulose membrane (NitroBind; Osmonics, Minnetonka, MN) and unspecific binding was blocked with tris(hydroxymethyl)aminomethane (Tris)-buffered saline containing 10% non-fat milk powder and 0.1% Tween – 20. A rabbit polyclonal antibody against mouse tPA (Molecular Innovations, Southfield, MI) served as the primary antibody. The secondary anti-rabbit antibody was conjugated with horseradish peroxidase (Jackson Immunoresearch Laboratories, West Grove, PA), and the antigen-antibody complex was detected with ECL Plus (Amersham Biosciences, Piscataway, NJ).

The plasminogen activator activity of anti-GPA scFv/PA and PA was confirmed using casein zymography. Aliquots of the SFM from anti-GPA svFv/PA- and PA-expressing cells were mixed with non-reducing Tris-glycine SDS sample buffer for zymography. The samples were resolved under non-reducing SDS-denaturing conditions on a 7.5% gel cast with 1% non-fat dry milk and 20μg/ml plasminogen incorporated into the gel before polymerization to detect PA activity (Wang et al-Deng, 2001). Thereafter, the gels were re-natured in Novex Zymogram Renaturating buffer (Invitrogen), and developed in Novex Zymogram Developing buffer (Invitrogen) per the manufacturer. EDTA (5 mM) was added to both buffers to block potential
metalloproteinase activity. Gels were stained with GelCode Blue stain (Pierce). Gels run in parallel in the absence of added plasminogen served as a control for PA activity (not shown).

The specificity of anti-GPA scFv/PA binding was confirmed using an immunocapture assay and WB. Briefly, mouse and human RBC ghost membranes were prepared (Schwartz et al., 1997) incubated with SFM from the induced S2 cells expressing anti-GPA scFv/PA for 1 h, then washed 3 times with PBS and lysed in a sample buffer (Invitrogen). The resultant RBC ghost lysates (equaled for total protein) were separated on 4-12% SDS-PAGE under non-reducing conditions. WB analysis of the samples to detect RBC ghost captured anti-GPA scFv/PA was performed as described above to detect scFv/PA in SFM medium. An aliquot of the SFM medium from the induced S2 cells expressing anti-GPA svFv/PA served as a positive control for the detection of fusion protein.

**Binding of anti-GPA scFv/PA to RBC.** We measured the binding of $^{125}$I-anti-GPA scFv/PA to mouse (target cells) vs human (negative control) RBC as previously described for an anti-CR1 mAb/tPA conjugate (Zaitsev et al., 2006). Briefly, RBC were washed by centrifugation (1200g) with PBS/3%BSA, resuspended in the same buffer to a hematocrit of 1% or 10% and incubated with various concentrations of $^{125}$I-scFv-PA for 1 h at 37°C (loading) with gentle rotation. Unbound reagent was eliminated by washing the RBC four times with a 20-fold volume of PBS-BSA. The residual radioactivity in the RBC pellets was measured in a γ-counter (Perkin Elmer). To determine RBC binding, mouse blood was collected in heparin and RBCs were prepared. $^{125}$I-anti-GPA scFv/PA was added to whole blood or washed RBC at a 50% hematocrit to final concentration of 40 μg/ml and binding was measured as described above.

**In vitro fibrinolysis.** The fibrinolytic activity of anti-GPA scFv/PA bound to RBC was measured using $^{125}$I-labeled fibrin clots, as described (Murciano et al., 2003). RBCs were incubated either with SFM medium from the induced S2 cells expressing anti-GPA svFv/PA or with purified anti-GPA scFv/PA for 1 h, washed 3 times with PBS and added to a solution
containing (6 mg/ml in PBS) trace labeled with $^{125}$I-fibrinogen. Clotting was induced by adding CaCl$_2$ and thrombin (20 mM and 0.2 units/ml final concentrations, respectively). The clots were then overlaid with 200 µl PBS, incubated at 37°C, and the radioactivity in the supernatants was measured in a γ-counter (Perkin Elmer).

In vivo tracing of RBC and RBC-anchored anti-GPA svFv/PA. Experiments were conducted under protocols approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Pennsylvania. Washed RBCs obtained from fresh anticoagulated mouse blood were labeled with $^{51}$Cr, as described previously (Zaitsev et al., 2006). $^{51}$Cr-RBC or $^{51}$Cr-RBC pre-loaded with $^{125}$I-anti-GPA scFv/PA was injected into anesthetized mice via the jugular vein. At designated times, aliquots of blood were drawn in heparin, the animals were sacrificed, and the radioactivity in the blood and major organs was measured.

Pharmacokinetic analysis of anti-GPA scFv/PA and PA. Adult C57BL/6 mice (Jackson Laboratories, Bar Harbor, ME) were studied. $^{125}$I-anti-GPA scFv/PA or $^{125}$I-PA (3-5 µg) was injected into anesthetized mice via the jugular vein. At the pre-designated times, 100 µl to 200 µl of blood was withdrawn in heparin, centrifuged at 1200g, and the radioactivity in the plasma and pellets was measured. The animals were sacrificed and the radioactivity in the organs was measured. In a separate set of experiments, RBCs obtained from mice were washed, labeled with $^{51}$Cr, loaded with $^{125}$I-anti-GPA scFv/PA at a dose of 20,000 molecules/RBC, and injected into anesthetized mice. Blood samples were collected and the major organs were harvested and analyzed for radioactivity as described for $^{125}$Iodine.

Fibrinolytic activity of anti-GPA scFv/PA loaded in vivo on circulating carrier RBC. We followed the protocol utilized previously for an anti-CR1 tPA conjugate (Zaitsev et al., 2006). Briefly, equimolar doses of anti-GPA scFv/PA or PA, providing an initial blood concentration of 0.85 µM (2mg/kg of PA and 4mg/kg of anti-GPA scFv/PA), were injected in 200 µl of saline vehicle into anesthetized mice via the jugular vein (drug-free saline was injected as a placebo.
control). Forty five min later, 100 μl aliquots of blood were drawn in the absence of anticoagulant, mixed rapidly with trace amounts of 125I-fibrinogen and allowed to clot in borosilicate tubes at 20°C. After 20 min maturation, clots were overlaid with saline and incubated at 37°C and the release of 125I was measured in a γ-counter (Perkin Elmer).

Effect of anti-GPA scFv/PA in a mouse model of carotid artery thrombosis. To test the anti-thrombotic potential of anti-GPA scFv/PA, we used a mouse model of acute severe carotid thrombotic occlusion in response to vascular injury inflicted by the adventitial application of FeCl₃ 30 min after administration of anti-GPA scFv/PA, PA or saline, as described previously (Murciano et al., 2003). An equimolar dose (0.85 μM) of anti-GPA scFv/PA or PA was injected into anesthetized mice as described above. Thrombosis was induced in the exposed contralateral carotid artery by applying a 1x2 mm piece of filter paper (Whatman No1) saturated with 15% FeCl₃ to the adventitia for 2 min. Time to occlusion of the vessel and total blood flow over the ensuing 30 min were measured by Doppler ultrasound using a 0.5VB flow probe connected to a recording system (Transonic Systems, Ithaca, NY).

Effect of anti-GPA scFv/PA in a mouse model of jugular vein thrombosis. To test the anti-thrombotic potential of anti-GPA scFv/PA, we used a mouse model of acute severe jugular vein thrombotic occlusion in response to vascular injury inflicted by the adventitial application of FeCl₃ 30 min after administration of anti-GPA scFv/PA, PA or saline, as described above (Murciano et al., 2003). An equimolar dose (0.85 μM) of anti-GPA scFv/PA or PA was injected into anesthetized mice as described above. Thrombosis was induced in the exposed contralateral jugular vein using 15% FeCl₃ for 2 min and the time to 50% occlusion and total blood flow maintained over the 30 min study were measured as described above.

Data analysis. All data are presented as the means plus or minus standard error of the means (SEM) of at least 3 separate experiments. Differences between groups were tested for
statistical significance using Student $t$ test or analysis of variance (ANOVA). Statistical significance was set at $P$ less than 0.05.
RESULTS

Design, synthesis and biochemical properties of the anti-GPA scFv/PA fusion protein. cDNA encoding scFv Ter-119 directed to mouse GPA was ligated to cDNA encoding the kringle II and protease domains of mouse tPA containing an additional Tenectaplace-type mutation using a (S4G)2A3 linker within the plasmid pMT-Bip His A (Fig. 1A). Transfection of this plasmid in S2 Drosophila cells lead to the expression of the fusion protein anti-GPA scFv/PA (scFv/PA). The PA part of the fusion protein was also cloned as a BglII/Xhol fragment into pMT-Bip His A and also expressed in S2 Drosophila cells. Non-reducing Western blot analysis with an antibody to mouse tPA was then performed to identify the two recombinant proteins. Anti-GPA scFv/PA and PA migrated as single bands with the predicted molecular weights of ~68 kDa and ~40 kDa, respectively) (Fig. 1B). Electrophoretic zymography demonstrated comparable PA activity for both proteins (Fig. 1C). To test the RBC binding capacity of anti-GPA scFv/PA, the medium from induced S2 cells transfected with the plasmid encoding anti-GPA scFv/PA was incubated with either mouse or human RBC membranes, washed and analyzed by immunoprecipitation, size fractionation and Western blotting as above. This analysis showed that anti-GPA scFv/PA bound to mouse, but not to human RBCs, which served in subsequent studies as the negative control (Fig. 1D).

RBC-binding properties of anti-GPA scFv/PA fusion. Both anti-GPA scFv/PA and PA were purified from cell media by affinity chromatography using ETI Sepharose to at least 95% purity confirmed by SDS-PAGE (data not shown). Purified 125I-labeled anti-GPA scFv/PA bound to mouse RBC membranes in a dose-dependent manner (Fig. 2A), whereas no significant binding was observed to human RBC at highest concentration tested (less than 100 copies bound per RBC, data not shown). Even at a 1% hematocrit, binding of anti-GPA scFv/PA (3x10^4 molecules per RBC) did not approach saturation, in agreement with the expression of 1x10^6 copies of GPA per RBC (Kina et al., 2000; Spitzer et al., 2004). The non-targeted PA moiety of the fusion did not bind to mouse RBC (open circle in Fig. 2B). Binding of anti-GPA
scFv/PA to mouse RBCs was rapid, reaching 50% of maximum within 10 minutes in whole blood (Fig. 2C).

**Fibrinolytic activity of RBC-bound anti-GPA scFv/PA.** To estimate whether RBC-bound anti-GPA scFv/PA activates plasminogen, mouse and human RBCs were incubated with medium from induced S2 cells transfected with anti-GPA scFv/PA encoding plasmid. Equal amounts of mouse or human RBCs pre-incubated with anti-GPA scFv/PA were washed and then added to a solution containing $^{125}$I-labeled fibrinogen and trace amounts of plasminogen prior to clotting by adding thrombin. Clot lysis was monitored by release of $^{125}$I-labeled fibrin degradation products into the supernatants. Mouse but not human RBCs pre-incubated with anti-GPA scFv/PA caused nearly complete lysis of the fibrin clot within one hour (Fig. 3A).

To analyze the fibrinolytic potency of RBC-bound anti-GPA scFv/PA, mouse or human RBCs were incubated with purified anti-GPA scFv/PA at a concentration chosen to attach ~20,000 molecules per RBC (Fig. 2). After removing unbound anti-GPA scFv/PA by washing, 2, 5 or 10 µl aliquots of the RBC preparations (50% hematocrit) were added to a solution containing $^{125}$I-labeled fibrinogen and trace amounts of plasminogen prior to clotting by thrombin, thereby achieving resultant concentrations of 1.5 nM, 3.75 nM and 7.5 nM anti-GPA scFv/PA in the clots, respectively. $^{125}$I-labeled degradation products were released into the supernatants in a dose- and time-dependent manner as a result of fibrinolysis (Fig. 3B). Release of radioactivity was not detected when mouse RBCs were replaced by human RBCs that do not bind anti-GPA scFv/PA (closed squares). Anti-GPA scFv/PA-RBC retain ~100% of their initial fibrinolytic activity at 24 hours and ~80% at 48 hours (data not shown).

We then used the same approach to compare the lysis of fibrin clots by equal amounts of free or RBC-bound anti-GPA scFv/PA. In samples incubated with free anti-GPA scFv/PA, an equal number of human RBC was added to account for the effect of cell incorporation on clot structure. Fibrinolysis was assessed after a 30 min incubation, the optimal time to reveal dose-
dependent differences (Fig. 3B). At rate-limiting doses, free anti-GPA scFv/PA caused more profound fibrinolysis than RBC-bound anti-GPA scFv/PA in vitro, likely due to diffusional limitations imposed by the RBC carrier (Fig. 3C).

**Binding of anti-GPA scFv/PA does not affect RBC biocompatibility and prolongs anti-GPA scFv/PA circulation time in mice.** To test the effect of anti-GPA scFv/PA binding on RBC biocompatibility and survival, washed $^{51}$Cr-labeled mouse RBCs, either intact or coated with $^{125}$I-labeled anti-GPA scFv/PA at a level of ~30,000 molecules per RBC, were injected into recipient mice. The amount of $^{51}$Cr in the blood and major organs was nearly identical in mice injected with either naive RBCs or anti-GPA scFv/PA coated RBCs at one and three hours post-injection (Figs. 4A and B). Of note, there was no detectable elevation in the uptake of RBCs in the lungs and spleen, indicating that anchoring of anti-GPA scFv/PA did not cause significant damage to the RBCs, nor cause them to aggregate in a manner that would enhance splenic clearance or induce mechanical retention in capillary-rich organs such as the lungs. More than 80% of injected $^{125}$I-anti-GPA scFv/PA remained in the circulation, with the rest found primarily in the liver (Figs. 4A and B). Centrifugation of heparinized blood samples showed that nearly 100% of both isotopes were recovered in the RBC pellet, consistent with stable binding of anti-GPA scFv/PA to RBC circulating in bloodstream as measured in vivo, with minimal elution into the plasma (Fig. 4A and B, insets).

Next, we compared the pharmacokinetics of $^{125}$I-anti-GPA scFv/PA vs $^{125}$I-PA injected directly into the bloodstream, simulating their clinical use. This setting differs from the “pre-loading” strategy described above, because here the binding occurs in the recipient’s bloodstream. Therefore, based on the binding kinetics showing complete binding by one hour after incubation of anti-GPA scFv/PA with RBC in vitro (Fig. 2C), tissue clearance within the first 30 min post-injection has the potential to eliminate a significant fraction of free drug. Nevertheless, at this time point there was a four-fold higher concentration of the anti-GPA scFv/PA fusion protein than non-targeted PA in the blood (Fig. 5A), and approximately 90% of
anti-GPA scFv/PA recovered in blood samples was associated with the RBC pellet (Fig. 5A, inset). Non-targeted PA and the unbound fraction of anti-GPA scFv/PA were taken up primarily by the liver; no appreciable accumulation of anti-GPA scFv/PA or PA was observed in the lungs or other major organs (Fig. 5A). It’s worth to mention, that when we measured the pharmacokinetics of free PA, we recovered ~40% of the injected dose in the major organs one hour post-injection. The half-life of free PA in circulation is several minutes due to rapid elimination by the liver and kidneys. Thus, we infer that the rest of the protein was excreted into urine and bile which could not collect and analyze as we did not use metabolic cages. The blood level of anti-GPA scFv/PA decreased gradually over time following a two-phase kinetics with a $T_{1/2}$ of approximately 2 and 10 hours for the rapid and slow phases, respectively (Fig. 5B). The major fraction of anti-GPA scFv/PA recovered in blood circulates bound to the RBCs for at least the first day after i.v. injection, while the plasma concentration peaked at 3-6 hours and was nearly undetectable by 24 hours (Fig. 5B, inset).

**Fibrinolytic activity of and prophylactic thrombolysis by anti-GPA scFv/PA injected in animals.** To test whether anti-GPA scFv/PA injected in mice retains its fibrinolytic activity, we analyzed the *ex vivo* lysis of clots formed from blood collected from mice without anticoagulants 45 minutes after i.v. injection of vehicle control (PBS), non-targeted PA or anti-GPA scFv/PA at equimolar doses. Clots formed from blood of animals injected with non-targeted PA did not undergo greater lysis than those formed from the blood of PBS-injected mice (Fig. 6A). In agreement with the higher blood level of RBC-targeted anti-GPA scFv/PA observed at this time point, we observed nearly complete lysis of clots formed from blood of mice injected with the fusion protein. This data indicate that the circulating, RBC-bound anti-GPA scFv/PA retains PA activity *in vivo*.

Based on this encouraging outcome, we then injected the same formulations in mice 30 minutes before inducing a thrombus in the carotid artery. Doppler analysis of perfusion through the carotid artery revealed that prophylactic administration of anti-GPA scFv/PA, but not non-
targeted PA, caused a three-fold delay in the time to vascular occlusion (Fig. 6B). Therefore, \textit{in vivo} loading of anti-GPA scFv/PA onto circulating RBCs provides thromboprophylaxis not feasible with its soluble non-targeted counterpart in this model, which is characterized by precipitous activation of platelets and coagulation cascade in a high-shear stress vessel.

Lastly, we tested anti-GPA scFv/PA in a model of jugular venous thrombosis. Venous ("red") clots, comprised predominantly of fibrin and RBC, might be especially amenable to RBC-tPA, compared to arterial ("white") thrombi, which are populated predominantly by platelets. Anti-GPA scFv/PA injected 30 min prior to jugular injury almost completely prevented vascular occlusion for the entirety of the experiment (Fig. 7 C and D), most likely due to expedited lysis of nascent "red" clots as they were forming in response to vascular injury (compare Fig. 7A and C, showing typical records of blood perfusion in the vessel). In contrast, untargeted PA delayed, but failed to prevent, vascular occlusion (Fig. 7B and D). Therefore, RBC-targeted anti-GPA scFv/PA, but not untargeted PA, preserved blood flow in the injured vein (Fig. 7E).
DISCUSSION

Thrombosis is the leading cause of mortality and disability in the United States (Jackson and Clagett, 1998). Thrombi are prone to recur within hours to days after a myocardial infarction, stroke, transient ischemic attack or pulmonary embolism, and when patients are immobilized (Wartenberg et al., 2004). Thromboembolism is also a common and dangerous complication of surgery, a setting that is especially difficult to manage due to the risk of exacerbating bleeding at the operative site. Therefore, situations in which patients are at high and predictable risk for thrombosis to occur (or recur), i.e., post-surgical patients, are known. However, anti-platelet and anticoagulant agents provide only limited prophylaxis and often pose considerable risk of bleeding, especially perioperatively (Zlokovic, 1997; Konstantopoulos and Mousa, 2001). Plasminogen activators (PA) are used for acute therapy of thrombosis in very circumscribed high risk settings (Topol et al., 1987; Holvoet et al., 1993). However, inadequate delivery (blood clearance within <15 minutes (Narita et al., 1995), inactivation by plasma inhibitors such as PAI-1 (Reilly et al., 1991) and impermeability of occlusive clots (Rijken et al., 2004) restrict the effectiveness of therapeutic fibrinolysis by PA. For example, very high doses of PAs (e.g., ~100 mg of tissue type PA, tPA) are needed to overcome these obstacles. At these high concentrations, drug may diffuse into hemostatic mural clots, predisposing to unwanted bleeding episodes, or into tissues such as the CNS (Wang et al., 1998) where it may cause cerebral hemorrhage, damage the blood-brain-barrier and direct neurotoxicity (Lo et al., 2003). Attempts to improve PA delivery and benefit/risk ratios have not yielded decisively better outcomes because the fundamental limitations with their use have not been overcome (Runge et al., 1996).

We hypothesized that prophylactic delivery of a PA would result in its incorporation into the interior of early nascent thrombi, arresting clot propagation and promoting clot lysis. This strategy would result in a more homogeneous drug delivery into the nascent thrombus rather than “therapeutic” fibrinolysis by external PA only effective on the clot surface. This, in turn,
would minimize the incidence of secondary emboli and thrombus re-formation. Short-term prophylactic use of PA in patients at high risk of imminent primary or recurrent thrombosis would also be predicted to reduce formation of occlusive clots impervious to delayed fibrinolysis.

Existing fibrinolytics are not used for prophylaxis due to their rapid clearance and serious side effects. Even newly designed mutant PAs with enhanced potency, including tPA variants with a mutated PAI-1 binding site and deleted accessory domains implicated in tPA clearance and adverse vascular signaling, i.e., Retavase and Tenekteplase, (Chapman et al., 2001) are likely to show limited diffusion into occlusive clots. Furthermore, all existing PAs are short-lived (<30 min), which makes them fundamentally inadequate for prophylaxis, and small (proteins with MW 30-60 kD, <10 nanometers diameter), which permits diffusion into hemostatic clots, increasing the risk of bleeding and increasing the propensity for collateral tissue damage, especially in the CNS. As of today, no PA has been designed for use as thromboprophylaxis.

Diverse drug delivery systems including liposomes have been employed to improve pharmacokinetics of plasminogen activators (Gupta et al., 2005; Elbayoumi and Torchilin, 2008). Previous studies from our group indicate that RBC provides a good carrier for fibrinolytics. Prior experiments in animal models of thrombosis documented that ex vivo coupling of tPA to carrier RBCs (RBC/tPA) provides effective and safe thromboprophylaxis, with the potential to shift the current paradigm for clot prevention (Murciano et al., 2003; Zaitsev et al., 2006; Danielyan et al., 2008). Such an ex vivo approach to RBC coating by drugs might be suitable in settings where transfusion is common, but would be less practical in other settings.

To avoid the need to couple tPA to isolated RBCs ex vivo followed by transfusion, we have targeted PA to circulating RBCs directly by conjugating tPA to a CR1 monoclonal antibody. We showed that the anti-CR1/tPA conjugate binds without harm to circulating RBCs in mice and provides safe and effective thromboprophylaxis (Zaitsev et al., 2006) comparable to that provided by infusion of RBC/tPA (Murciano et al., 2003).
However, antibody conjugates, useful in animal studies, are sub-optimal for clinical use and would be challenging to produce in sufficient quantities. Furthermore, there is the potential to form large (MW>270 kDa), heterogeneous and multimeric conjugates that may activate cellular defense mechanisms and complement via Fc-fragments. To circumvent these problems and to achieve predictable coating levels over a wide range of drug concentrations, we designed recombinant PA variants fused to an antigen-binding vehicle (single chain variable fragment [scFv]) directed to mouse glycophorin A (Spitzer et al., 2004).

Expression of scFv/PA through recombinant technology enables large-scale production of homogeneous monovalent scFv/PA fusion proteins (Holvoet et al., 1993). As scFvs lack the Fc portion of an intact antibody, the risk of immune-mediated side effects lessened. Established techniques for humanization and methods to reduce the potential immunogenicity of scFv chimeras further help to minimize the potential for eliciting immune reactions (Almagro and Fransson, 2008). Lastly, the modular recombinant format used in our studies supports the synthesis of targeted variant PA pro-drugs lacking domains that may elicit untoward effects.

The results shown in the present paper provide proof-of-principle for the proposed strategy. Anti-GPA scFv/PA (Fig. 1) bound to RBCs and invested them with PA activity (Fig. 2 and 3). When injected at equimolar doses in mice, RBC-targeted anti-GPA scFv/PA exhibited markedly higher fibrinolytic activity in the circulation than a non-targeted PA variant (Fig. 6 and 7). This was likely due to binding of the anti-GPA scFv/PA to circulating RBCs, which enhanced its circulation time and the bioavailability in vivo (Fig. 5). This major change in the pharmacokinetics of anti-GPA scFv/PA compared to non-targeted PA overrode a partial reduction of its resultant specific fibrinolytic activity in vitro compared to free anti-GPA scFv/PA (Fig. 3C). This inequity of fibrinolytic potency in vitro may reflect restricted diffusional freedom within the clot of anti-GPA scFv/PA bound to the large RBC carrier, which is relatively immobile in the clot meshwork. Therefore, it is likely that in vivo comparison of fibrinolytic activity of anti-
GPA scFv/PA vs PA at equipotent rather than equimolar doses would reveal an even more profound advantage of the RBC-targeted anti-GPA scFv/PA over its non-targeted counterpart.

Testing of anti-GPA scFv/PA in mouse models of thrombosis provided important new in vivo findings relevant to pharmacological properties of this potential biotherapeutic. Anti-GPA scFv/PA dissolved venous vs arterial thrombi more effectively, as expected. However, alleviation of arterial thrombotic occlusion by anti-GPA scFv/PA implies that sufficient drug-loaded RBC’s were incorporated into the clot even at high shear to achieve their intended effect.

The results reported here warrant further systematic evaluation of dosing, timing and duration of beneficial and potential adverse effects of anti-GPA scFv/PA in laboratory animals. For example, the duration of prophylaxis here was limited and would need to be extended to be of clinical use. However, the stability of the RBC-scFv/PA complex in vivo portends protracted thromboprophylactic activity although this remains to be proven. Moreover, subtle effects on GPA function with protracted use not evident in this study will have to be excluded. It will also be necessary to replace the anti-murine GPA scFv with one against human GPA to invest it with clinical utility. On the other hand, this study provides a modular template to design diverse iterations of this novel class of biotherapeutics, utilizing recombinant and mutagenesis techniques to vary their affinity and binding sites on RBC, and molecular structure of plasminogen activator domains involved in regulation such important auxiliary features as resistance to plasma inhibitors, interaction with vascular receptors and regulation of the enzymatic activity by fibrin and other components of clots and clotting cascade. These additional molecular modifications may further enhance the clinical utility, safety, potency and specificity of RBC-targeted plasminogen activators.
REFERENCES


Footnotes

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Legends for Figures

Figure 1. Molecular design, expression and characterization of anti-GPA scFv/PA fusion protein. (A) Schematic diagram describing the cloning strategy for the fusion construct anti-GPA scFv/PA. Variable domains of the heavy chain and light chains of the antibody were linked by a \((\text{Gly}_4\text{Ser})_3\) linker and then fused to the N-terminus of the kringle II/protease domain fragment of mouse tPA by a \((\text{Ser}_4\text{Gly})_2\text{Ala}_3\) linker. The completed construct was then cloned into the SpeI and XhoI sites in the pMT/BIP/V5/HisA expression vector. The DNA fragment encoding kringle II and protease domain of mouse tPA was PCR amplified and cloned into BglII amd XhoI sites in the same vector. (B) Western blot analysis of 40 µL culture medium from S2 cells expressing either anti-GPA scFv/PA fusion protein or PA after induction by 0.5 mM CuSO\(_4\). (C) Casein zymography analysis of culture medium from S2 cells expressing either the anti-GPA scFv/PA fusion protein or the PA after induction by 0.5 mM CuSO\(_4\). (D) Western blot analysis of protein immunocaptured with mouse (m) or human (h) RBC ghosts: lane 1 = lysate of hRBC ghosts incubated in culture medium from S2 cells expressing anti-GPA scFv/PA fusion protein; lane 2 = lysate of mRBC ghosts incubated in culture medium from S2 cells expressing anti-GPA scFv/PAfusion protein; lane 3 = lysate of hRBC ghosts; lane 4 = lysate of mRBC ghosts; lane 5 = 40 µl of culture medium from S2 cells expressing anti-GPA scFv/PA fusion protein (positive control).

Figure 2. Specific binding and binding kinetics of anti-GPA scFv/PA fusion protein to RBC expressing mouse GPA. (A) Dose dependent binding of \(^{125}\text{I}\)-anti-GPA scFv/PA fusion protein to washed mouse RBC at 10% hematocrit RBC suspension determined
after elimination of unbound material (n=3). (B) Dose dependent binding of $^{125}$I-anti-GPA scFv/PA fusion protein (closed circles) vs PA (open circles) to washed mRBC at 1% hematocrit RBC suspension determined after elimination of unbound material (n=3). (C) Binding kinetics of $^{125}$I-anti-GPA scFv/PA fusion protein to mRBC in whole heparinized blood vs the same hematocrit washed mouse RBC suspension in PBS/BSA buffer (each time point n=3). Mean ± 1 standard error from the mean (SEM) are shown.

**Figure 3. Fibrinolytic activity *in vitro* of RBC-bound and free anti-GPA scFv/PA fusion protein.** (A) Fibrinolytic activity of mouse or human RBC (mRBC vs hRBC) pre-incubated for 1 hour in the medium of the S2 cells expressing anti-GPA scFv/PA fusion protein. RBCs were then washed and 20 µl of 50% RBC suspension was incorporated in a forming 200 µl fibrin clot. After maturation fibrin clots were brought to 37°C and clot lysis was monitored (n=3). (B) Dose dependent lysis of the fibrin clot by mouse RBC loaded with anti-GPA scFv/PA fusion protein. Incorporation in a fibrin clot of 2, 5 and 10 µl of loaded mouse RBC, 50% hematocrit suspension, correspond to 1.5, 3.75 and 7.5 nM of anti-GPA scFv/PA in a clot, respectively (n=3). 10 µl of human RBC suspension incubated with anti-GPA scFv/PA at the same conditions as mouse RBC and 10 µl intact mouse RBC suspension incorporated in the clot served as controls. (C) Comparison of fibrinolytic activity of anti-GPA scFv/PA in a free and RBC-bound state. Equal amounts of anti-GPA scFv/PA were incorporated in a forming fibrin clots in mouse RBC bound or free state (the structure of the clot was kept the same by incorporation of an equal number of human RBC). Clot lysis was monitored for 30 minutes (n=3). Dash line shows the spontaneous lysis of fibrin clots with incorporated intact RBCs. In each experiment the mean ± 1 SEM are shown.
Figure 4. Anti-GPA scFv/PA loading does not damage carrier RBC. $^{51}$Cr-labeled mouse RBC, either naïve or loaded at the level of 20,000 /RBC with $^{125}$I-anti-GPA scFv/PA fusion protein, were injected into mice. The animals were killed 1 hour (A) and 3 hours (B) afterwards, and the amount of $^{51}$Cr and $^{125}$I was measured in blood and the main organs (n=4). The insets in both panels indicate $^{51}$Cr and $^{125}$I distribution in blood components at the indicated times. In each experiment the mean ± 1 SEM are shown.

Figure 5. Pharmacokinetics of $^{125}$I-anti-GPA scFv/PA fusion protein vs $^{125}$I-PA in mice. (A) Organ distribution of $^{125}$I-anti-GPA scFv/PA fusion protein vs $^{125}$I-PA after 1h circulation in mice. Inset: blood components distribution of $^{125}$I-anti-GPA scFv/PA fusion protein vs $^{125}$I-PA after 1h circulation in mice. (B) Blood clearance of $^{125}$I-anti-GPA scFv/PA fusion protein during 24h circulation after IV injection in mice. Inset: percent of $^{125}$I-anti-GPA scFv/PA fusion protein recovered in plasma vs RBC pellet in blood obtained at the indicated times after IV injection of $^{125}$I-anti-GPA scFv/PA fusion protein in mice. (The number of animals in all experiments is 5 per group). In each experiment the mean ± 1 SEM is shown.

Figure 6. Fibrinolytic activity of circulating anti-GPA scFv/PA/mRBC complex in arterial thrombosis. (A) Fibrinolytic activity recovered in mouse blood samples obtained 45 minutes after IV injection of 4 mg/kg dose of anti-GPA scFv/PA fusion protein, 2 mg/kg PA or saline. n=4 per group; p< 0.05. (B) Occlusive thrombi were formed in the carotid artery of mice by applying FeCl₃ to adventitia. Thirty minutes before injury 4 mg/kg dose of anti-GPA scFv/PA fusion protein, 2 mg/kg PA or saline we injected IV (jugular vein). Time of complete vessel
occlusion was determined with Doppler Ultrasound. Data shown as mean ± SEM, n=8 per group, p< 0.05.

**Figure 7. Fibrinolytic activity of circulating anti-GPA scFv/PA/mRBC complex in venous thrombosis.** Panels A, B, and C show the typical records of the blood flow in injured jugular vein in mice monitored by Doppler. Saline (A), untargeted tPA (B), or equimolar dose of anti-GPA scFv/PA fusion protein (C), was injected IV in a contralateral jugular vein thirty minutes before injury inflicted by FeCl₃. Panel D shows analysis of data collected in these animal groups depicted as the time needed to attain the 50% reduction of the blood flow after the injury. Panel E shows analysis of the data collected in these groups presented as percent of retention of the blood flow in the jugular vein during 30 min after induction of injury in mice treated with prophylactic administration of equimolar doses of PA vs anti-GPA scFv/PA 30 min before injury. In panels D and E, the mean ± SEM is shown, n=6 per group, p< 0.05.
Fig 4
**Fig 5**

**A**

- **Graph A**: Tissue level, % ID
- **Y-axis**: Tissue level, % ID
- **X-axis**: blood, lung, liver, kidney, spleen, heart
- **Legend**:
  - anti-GPA scFv/PA
  - PA

**B**

- **Graph B**: Blood level, % ID
- **Y-axis**: Blood level, % ID
- **X-axis**: Time, hours (0.5, 1.0, 3.0, 6.0, 24.0)
- **Legend**:
  - RBC
  - plasma

**Inset Bar Graph**

- **Y-axis**: RBC/plasma ratio, %
- **X-axis**: anti-GPA scFv/PA, PA
Fig 6

(A) Blood clot lysis, %
- PA
- anti-GPA scFv/PA
- PBS

P < 0.05

(B) Time to occlusion, min
- PA
- anti-GPA scFv/PA
- PBS

P < 0.05