Advanced drug delivery systems for anti-thrombotic agents

Colin Greineder,¹,³ Melissa D. Howard,¹ Ron Carnemolla,¹ Douglas B. Cines,² and Vladimir Muzykantov¹*

¹Department of Pharmacology and Center for Targeted Therapeutics and Translational Nanomedicine, Perelman School of Medicine, University of Pennsylvania;
²Department of Pathology and Laboratory Medicine, Perelman School of Medicine, University of Pennsylvania
³Department of Emergency Medicine, Perelman School of Medicine, University of Pennsylvania

*Correspondence: Vladimir Muzykantov, University of Pennsylvania, Perelman School of Medicine, Department of Pharmacology and Center for Targeted Therapeutics and Translational Nanomedicine, TRC10-125, 3600 Civic Center Boulevard, Philadelphia, PA 19104, USA. Tel +1-215-898-9823; fax +1-215-573-9135; muzykant@mail.med.upenn.edu

First two authors equally contributed to the paper.

Running title: Drug Delivery Systems for Anti-thrombotic Agents
Abstract:

Despite continued achievements in anti-thrombotic pharmacotherapy, difficulties remain in managing patients at high risk for both thrombosis and hemorrhage. Utility of anti-thrombotic agents (ATAs) in these settings is restricted by inadequate pharmacokinetics and narrow therapeutic indices. Use of advanced drug delivery systems (ADDS) may help to circumvent these problems. Various nanocarriers, affinity ligands, and polymer coatings provide ADDS that have the potential to help optimize ATA pharmacokinetics, target drug delivery to sites of thrombosis, and sense pathologic changes in the vascular microenvironment, such as altered hemodynamic forces, expression of inflammatory markers, and structural differences between mature hemostatic and growing pathological clots. Delivery of ATA using biomimetic synthetic carriers, host blood cells, and recombinant fusion proteins that are activated preferentially at sites of thrombus development has shown promising outcomes in preclinical models. Further development and translation of ADDS that spare hemostatic fibrin clots hold promise for extending the utility of ATAs in the management of acute thrombotic disorders through rapid, transient, and targeted thromboprophylaxis. If the potential benefit of this technology is to be realized, a systematic and concerted effort is required to develop clinical trials and translate the use of ADDS to the clinical arena.
Introduction

Thrombosis is a key event in the pathogenesis of myocardial infarction, venous thromboembolism (VTE), stroke and other ischemic conditions that comprise the most common and growing cause of morbidity and mortality in industrialized communities. Development of anti-thrombotic agents (ATAs) is a global medical priority. ATAs may be subdivided conceptually into anti-platelet agents, anticoagulants, and fibrinolytics. Therapeutics within each category differ with respect to mechanism of action, time to onset, duration of effect, route of administration, and ease of monitoring and reversal (Supplemental Table 1).

Traditionally, anti-platelet agents and anticoagulants have been perceived primarily as preventing clotting, whereas fibrinolytics have been viewed as 'rescue' therapy in acute settings (Supplemental Figure 1). Yet, many ATAs have dual preventive and therapeutic effects in acute settings due to the dynamic nature of clots, which may involve repetitive cycles of clot accretion, maturation, and dissolution. For example, anticoagulants are often used for treatment as well as prevention, due to their ability to both inhibit thrombus growth and to enhance endogenous fibrinolysis.

While many ATAs are in clinical use, there continues to be a need to optimize efficacy and safety, especially in patients at high risk of bleeding and thrombosis. Narrow therapeutic indices complicate management of patients with commonly encountered clinical scenarios, including the prevention of venous thrombosis in high-risk surgical patients, such as those with trauma or cancer (Figure 1). Current approaches to thromboprophylaxis in such patients, though successful, necessitate a careful and oft-times difficult balance between the estimated risk of thrombosis with drug interruption and the risk of bleeding due to premature resumption of thromboprophylaxis.\(^1\)\(^-\)\(^3\) Stroke and delayed onset cognitive dysfunction attributed to thromboemboli remain a serious and increasingly recognized late complication of surgery. The risk of causing or accentuating bleeding also constrains the use of plasminogen activators (PAs) in patients with sub-massive pulmonary embolism and thrombotic stroke. In comparison with recent successes in the development of novel anti-platelet agents and anticoagulants, there have been fewer advances in fibrinolytic therapy since the introduction of tPA. However, even partial preservation of perfusion affected by obstructing thrombi may suffice to prevent ischemic tissue injury. Therefore, it is not surprising that research in the use of advanced drug delivery systems (ADDS) has mainly focused on fibrinolytic therapy.

A fundamental limitation shared by all current ATAs is the inability to prevent or reverse thrombosis without interfering with hemostasis. ADDS have the potential to help overcome this limitation and thereby broaden the therapeutic window of anti-thrombotic interventions\(^4\) through two key, often intertwined functions: i) prolonging the circulation time and minimizing the undesirable drug/host interactions en route to the therapeutic site; and, ii) spatiotemporally
localizing drug action. In this article we briefly review using ADDS for anti-
thrombotic agents.

2. Prolongation of ATA half-life and intravascular activity. Multiple approaches have been devised to extend the half-life and activity of ATAs in the bloodstream, including polymer coating and the use of drug carriers. Most carriers are artificially assembled supramolecular structures composed of biocompatible elements, with sizes ranging from tens of nanometers to a few microns (Figure 2).

2.1. Stealth PEG coating, liposomes and polymeric carriers. The first approach to prolong the half-life of ATA involved conjugation with polyethylene glycol (PEG). PEG chains form a hydrated shell that enhances drug solubility and limits renal filtration, cellular uptake, proteolytic degradation, and immunogenicity (“stealth” ADDS, an approach that has proven effective for some biotherapeutics). PEGylation of tPA, streptokinase (SK), urokinase (uPA), and staphylokinase (Sak) has been tested extensively. PEG-PA conjugates prepared with an optimal extent of protein modification exhibit a prolonged circulation time and minimal loss of activity. In a phase IIa trial PEG-Sak had therapeutic efficacy similar to tPA, but the study revealed hemorrhagic complications.

Liposomes, spherical vesicles with diameters ~100-250nm composed of a phospholipid bilayer (that may contain PEGylated phospholipids) surrounding an aqueous core, are used clinically to deliver chemotherapeutic agents. PEG-liposomes protect ATAs from degradation, reduce antigenicity, and increase plasma half-life. ATAs encapsulated in PEG-liposomes showed encouraging results in vitro and, to a more limited extent, in animal models. For example, liposomal PAs provided superior reperfusion vs free PAs in rabbit models of jugular vein and carotid artery thrombosis.

Interactions with cells and lipoproteins, as well as the intense PEGylation needed for a strong “stealth” effect, destabilize liposomes, limiting their life-time in bloodstream. More stable carriers based on amphiphilic polymers (frequently PEG block copolymers) have also been tested to extend the circulation time of ATAs. Examples include chitosan/poly(lactic-co-glycolic acid) nanoparticles for tPA, glycol chitosan/PEG nanogels for uPA, as well as polyglycerol and poly(amido amine) dendrimers for SK. In a canine model of coronary artery thrombosis, SK encapsulated within PEGylated particles caused faster reperfusion than free SK. However, in vivo data for polymeric carriers are even more limited than for PEGylated and liposomal ATAs.

These approaches prolong the circulation of ATAs on par with recombinant tPA variants (T1/2 in blood ~5 min for tPA vs ~20-40 min for Reteplase and the more potent PAI-1 resistant Tenecteplase), which have not provided a decisive therapeutic advantage. Additional studies are needed to
determine the factors that contribute to clot resistance in vivo, including poor perfusion of occluded vessels, local high levels of PAI-1, clot retraction and high fibrin affinity limiting clot penetration, among others.\textsuperscript{26} Enlargement of ATAs by a carrier (e.g., from a few nanometers up to hundreds) further impedes clot permeation. The mechanism by which encapsulated ATAs could interact with fibrin and distinguish between occlusive vs hemostatic thrombi is unclear. Additionally, the prolongation of half-life is still not sufficient for prophylactic administration, which might enable ATAs to be delivered to the interior of nascent clots thereby overcoming permeability issues.

### 2.2. ATA carriers sensitive to ultrasound and hydrodynamic forces.

One approach toward circumventing these problems is to release ATA from carriers at the intended therapeutic site. Echogenic liposomes (ELIP), which contain both gas and fluid in the core, can be used for drug delivery and ultrasound imaging. Locally applied ultrasound facilitates reperfusion of thrombotic vessels in animals injected with tPA-loaded ELIPs,\textsuperscript{27,28} most likely by accelerating tPA release and disintegrating clots directly, enhancing drug permeation. PEG-gelatin nanocarriers also prolonged the circulation of tPA and released it in ultrasound-responsive fashion, improving recanalization of occlusive clots in rabbits\textsuperscript{29}. Further refinements of the ELIP approach are warranted, as potent ultrasound may inactivate PAs\textsuperscript{30} and cause vascular damage. Some clots may not be amenable to this approach due to unknown location or risk of hemorrhage.

To avoid the need for an external mechanism to release ATAs, “smart” carriers sensitive to hemodynamic changes have recently been designed (Supplemental Figure 2). These shear-activated “nanotherapeutics” (SA-NTs)\textsuperscript{31} represent micron-size (~4 μm diameter) aggregates of tPA-coated polymeric nanoparticles (~200 nm diameter) that are relatively stable under normal flow (10-30 dyne/cm\textsuperscript{2}) but dissociate into nanoparticles at the elevated shear stress levels encountered in areas of arterial stenosis (>100 dyne/cm\textsuperscript{2}). Nanoparticles experience lower drag forces than their parental micron particles and accumulate in affected vessels, thereby delivering the ATA. In a mouse model of thrombotic arterial stenosis, SA-NTs prolonged the time to vessel occlusion from ~10 to ~30 min in vivo and cleared pulmonary emboli in an ex vivo model at a 100-fold lower dose than free tPA. Lenticular lipid-based vesicles that show shear-stress induced drug release have also been developed but not yet evaluated in vivo.\textsuperscript{32}

Non-spherical carriers imitating the hemodynamic properties of platelets\textsuperscript{33} or red blood cells (RBCs)\textsuperscript{34,35} circulate in model microfluidic systems and animals for extended periods of time, exceeding the limits of traditional nanocarriers.\textsuperscript{34} Elongated PEG-polymeric filomicelles that align with blood flow and thereby avoid collisions with vascular cells have circulation times on the order of days as opposed to hours as seen with spherical PEG-coated carriers.\textsuperscript{36} This may allow them to integrate into growing thrombi. For example, when coated with a recombinant VWF-A1 fragment that targets the GPIb\textalpha region on activated
platelets, “synthetic platelets” were incorporated into clots. This is a new area of research, and the in vivo behavior and therapeutic efficacy of ATA loaded particles remain to be tested. Challenges of using such polymeric carriers include inactivation of enzymatic drugs during loading and the need for substrate permeability or controlled release. Encapsulation of active enzymes in filomicelles has been reported, which provides hope they can find utility for delivering ATAs. These studies also point toward exploring the natural carriers, blood cells, which inspired their development.

2.3. ATA delivery by erythrocytes: Transient Targeted Thromboprophylaxis (TTT). RBCs (~6 micron discs with a circulation time of ~120 days in humans) have many features of ideal carriers for drugs that act in the bloodstream, especially when sustained action is needed. Drugs can be encapsulated within isolated RBCs followed by resealing and injection. Promising results from transfusion of RBC-encapsulated drugs in neurological, oncological and other diseases have been reported. Biocompatible coupling of ATAs to the surface of carrier RBCs represents a preferable alternative to encapsulation.

Once clots are established, they become progressively impermeable to RBCs. While this feature negates the utility of RBC carriers for therapeutic thrombolysis, it also provides a key safety feature for TTT. In theory, incorporation of PAs within the interior of nascent thrombi will arrest clot propagation and dissolve clots from within more effectively than PAs permeating clot from the surface. If feasible, anticipatory delivery of PAs within nascent thrombi might prevent vascular occlusion in patients at imminent risk of thrombosis without lysing hemostatic clots. Although the inherent limitations of PAs discussed above preclude their use in thromboprophylaxis, coupling PAs to RBCs surmounts these problems (Figure 3).

RBC carriers were first devised for PAs. Studies in mice, rats and pigs have shown that conjugation of PAs to RBCs generates a biocompatible RBC/PA complex that does not lyse hemostatic clots formed as recently as 10 min post-surgery that are readily lysed by free PA. The circulation of RBC/PA in the blood is several orders of magnitude longer than PA (many hours vs minutes), which allows inclusion of prophylactically injected RBC/PA within nascent intravascular clots followed by rapid lysis and reperfusion, in settings where a 10-fold higher dose of soluble PA is ineffective. As clots mature, they generate less thrombin and undergo physical retraction, which markedly reduces penetration by RBCs and even by tPA. Moreover, hemodynamic forces divert circulating RBCs from mural clots. For these and other reasons, RBC/PA show selectivity towards lysing nascent clots forming in the lumen compared with preformed clots.

The RBC glycocalyx attenuates inactivation of coupled tPA by PAI-1, without affecting its fibrin affinity, which facilitates RBC/tPA entrapment in
Coupling urokinase to RBCs blocks its interaction with cellular uPA receptors and limits unwanted adhesion of RBC/PA to vessel walls. RBC/PA entrapped within growing clots rapidly forms cavities that merge under flow to create patent channels that permit expedited perfusion by RBCs prior to extensive disintegration of clot structure emanating from the clot surface by soluble PAs.

Intravenously injected RBC/PA rapidly lyses nascent cerebrovascular thrombi leading to reperfusion, protection of brain tissue and improved survival. In animal models of traumatic brain injury and intracranial hemorrhage, RBC/PA injected shortly after trauma attenuated secondary thrombosis and neuronal death seen in control animals. In contrast, soluble PA exacerbated CNS bleeding, neurotoxicity and lethality. Further, RBC/tPA preserved cerebral vascular autoregulation disrupted by cerebral hypoxia in contrast to tPA, which increased cerebrovascular contractility and tissue ischemia. Finally, tPA aggravated inflammatory pathways and brain injury in pig models of global cerebral hypoxia and focal thrombotic ischemia, whereas RBC/tPA activated anti-inflammatory pathway(s) and attenuated brain injury. Therefore, coupling to RBC effectively switches the PA signaling profile in the CNS from injurious to protective, permitting a reduction in dosing and mitigating ischemia by enhancing clot lysis, while reducing deleterious intravascular and extravascular signaling.

These “RBC loading/transfusion” studies show the potential advantages of prophylactic thrombolysis – a concept that might expand the current paradigm of fibrinolytic therapy.

2.4. Prolonging ATA circulation: challenges and perspectives. Artificial carriers designed to prolong half-life and improve delivery of ATAs are not yet ready for clinical use. Although each carrier has its own advantages and disadvantages (Table 1), none prolong the longevity of ATA in the circulation sufficiently to capitalize on the dynamic nature of developing clots. Their therapeutic utility for fibrinolysis is impeded by the impermeability of maturing clots. Flow-sensitive nanocarriers may overcome some of these limitations in settings characterized by partial thrombotic occlusion and pathologically high shear stress, although their utility in low shear conditions may be more limited.

RBCs markedly prolong the circulation and limit the side effects of fibrinolytics sufficiently in order to envision their utility in thromboprophylaxis. RBC/tPA prevented venous and arterial thrombotic occlusions in animal models. The relative effectiveness of RBC- vs platelet-ATAs in preventing venous vs arterial thrombosis requires further study. Clinical heterogeneity in clot maturation and the attendant risk of bleeding and clotting will require extensive preclinical investigation to determine dosing and timing to optimize the benefit to risk ratio of any intervention.

3. Intravascular targeting of ATA. A variety of affinity ligands including
antibodies and their single chain variable fragments (scFvs), clot components, aptamers and peptides have been employed to target fibrinolytics and, to a lesser extent, anticoagulants. The most intensively investigated targets for delivering ATAs have been clotting factors, components of clots, blood components that may deliver a cargo to clots and endothelial surface determinants providing anchoring of cargoes in the vascular lumen.

Chemical conjugation of large ATA molecules such as heparin and proteins (e.g., tPA) with these ligands generated products that vary in specific activity, molecular structure, valence and size, which impedes industrial and clinical translation. Recombinant constructs fusing the drug with scFv fragments (Supplemental Figure 3) are advantageous, because: i) Binding of monovalent scFvs does not cross-link epitopes that could lead to cellular agglutination, activation and endocytosis; ii) scFv fusions lack Fc-fragment mediated side effects; iii) Modular formats support rapid synthesis of iterations of ATA fusions bearing different cargoes; iv) Affinity of fusions can be optimized in vitro; v) Industrial GMP-production of homogeneous scFv fusions and techniques for scFv humanization minimizing the likelihood of immune reactions are established.

The recombinant format of ATA fusions makes it amenable to introducing functionally advantageous mutations into the cargo. For example, uPA is synthesized as a single chain zymogen (scuPA) in contrast to tPA, which is constitutively active. Proteolysis of scuPA between Lys and Lys yields an ~30 kD zymogen, low-molecular weight (lmw) scuPA that lacks the growth factor-like and kringle domains that bind to CD87/uPAR and other receptors on vascular cells and leukocytes, triggering potentially deleterious signaling pathways. Cleavage of scuPA or lmw scuPA by plasmin at Lys-Ile yields enzymatically active two-chain molecules, tcuPA and lmw tcuPA, which produce plasmin.

Insertion of cleavage sites for proteases involved in clotting (e.g., thrombin and Factor Xa) creates a gradient for activating latent ATA cargoes emanating from the site of incipient thrombosis (Figure 4 and Supplement Figure 3). For example, physiologic activation of scuPA by plasmin may not suffice to prevent thrombosis and tcuPA is rapidly inactivated by plasma inhibitors. Thrombin also inactivates uPA by cleaving Arg-Phe, negating its effect at sites of active thrombosis. These problems can be circumvented by deleting Phe and Lys to yield a plasmin-resistant mutant activated by thrombin (uPA-T). This prodrug is not activated by plasmin in vivo (thus avoiding systemic effects and premature inactivation), but engenders activation by thrombin locally at sites of nascent thrombosis within seconds of clotting. This mutant, scuPA-T, is thereby invested with a more favorable targeting profile.

3.1. Targeting ATA to thrombi. The discovery of fibrin-specific epitopes absent in fibrinogen provided the rationale and tools for targeting ATA to thrombi. Fibrin-targeted ATA conjugates, nanocarriers and scFv fusions with
thrombin-induced ATA activation and release (Figure 4) show greater anti-thrombotic therapy in animal models.\textsuperscript{71,72}

Platelets have also been exploited as carriers for ATA, especially for arterial clots.\textsuperscript{73} Ligands recognizing the conformational change in GPIIb/IIIa or P-selectin exposed upon platelet activation might direct ATA or ATA-loaded carriers to growing platelet-rich clots.\textsuperscript{74,75} For example, uPA conjugated to a monoclonal anti-GPIIb/IIIa scFv binds to platelets and platelet-rich thrombi and lysed arterial thrombi.\textsuperscript{76} Some of the conjugate’s activity might derive from blocking the fibrinogen receptor GPIIb/IIIa, like Abciximab\textsuperscript{77}. Platelet-targeted ATA may accumulate in thrombi by binding to: i) resting platelets in the circulation with subsequent incorporation into growing clots and/or ii) platelets within clots themselves (Figure 4C). The contribution of the latter pathway needs to be better defined, for example by tracing injected ATA derivatives in blood and thrombi.\textsuperscript{78}

However, the conundrum of targeting ATA to clot components is that their accessibility diminishes rapidly during thrombosis. Paradoxically, the higher the affinity of a ligand for fibrin, the greater is its retention on the clot surface and the less its penetration.\textsuperscript{79} Yet, clot-targeted ATAs have insufficient longevity for use as hemostrophosphylaxis. Finally, no existing ligand distinguishes pathological from hemostatic clots.

3.2. Targeting ATA to circulating RBCs. ATA targeting to RBCs, i.e., a logical extension of the principle described in Section 2.3, helps circumvent many of these issues and bypasses the complexity, delays and potential risks associated with ex vivo coupling to allogeneic RBC. As a prototype, tPA conjugated with a monoclonal antibody to RBC glycoprotein complement receptor 1 (CR1) bound rapidly to RBCs in the bloodstream and circulated safely for many hours after injection in mice, providing prophylactic thrombolysis without hemorrhagic side effects, similar to preformed RBC/tPA.\textsuperscript{80} Subsequently, a scFv/tPA fusion targeted to RBC glycophorin-A related antigen (GPA) was developed that bound safely to circulating RBC and had anti-thrombotic effects in mouse models of thrombosis qualitatively similarly to RBC/tPA and superior to tPA, with an additional advantage of wide range of dosing options (~10\textsuperscript{6} copies of GPA vs 2x10\textsuperscript{3} CR1 copies per RBC, respectively).\textsuperscript{80-82}

To enhance spatiotemporal control, the tPA cargo was replaced with lmw-scuPA in which the natural plasmin-sensitive activation site was replaced with a thrombin-sensitive site. Animal studies showed that the resultant scFv/uPA-T fusion: i) binds safely to RBCs after IV injection; ii) circulates as a RBC-bound prodrug; iii) is activated by thrombin; and, iv) provides effective and safe TTT with higher fidelity and efficacy than even scFv/tPA (Figure 5A).\textsuperscript{82}

Fibrinolytics are not the only potential ATA cargo suitable for RBC carriage. For example, recombinant soluble thrombomodulin (sTM), a natural anticoagulant that shows encouraging clinical results,\textsuperscript{83} has a half-life in the
circulation that can reach several days after subcutaneous injection. This can be extended many fold by targeting to RBCs, and the onset of its action can be expedited by intravenous administration of sTM fused with scFv anti-GPA. In animal models, RBC-targeted anti-GPA scFv/TM: i) binds safely to RBCs; ii) generates activated protein C (APC) from PC in the presence of thrombin; iii) has a half-life ~100-fold longer than sTM after IV injection; and, iv) prevents occlusive arterial and venous thrombi from forming at doses that are orders of magnitude lower than sTM. TM and APC have multifaceted anti-thrombotic and anti-inflammatory activities. RBC-targeted scFv/TM may find therapeutic and prophylactic utility in such intertwining pathologies (Figure 5B).

In summary, targeting to carrier RBCs might offer a preferred approach to deliver and locally activate anti-thrombotic prodrugs within incipient thrombi, while sparing even recent hemostatic clots that become impermeable to RBCs within minutes. Targeting scFv/uPA-T and scFv/TM to RBCs might provide TTT for patients at high risk of thrombosis in the early post-surgical period or with related conditions that combine acute risk of thrombosis with a heightened risk of hemorrhage.

3.3. Targeting ATA to endothelial cells. Endothelial cells (ECs) utilize natural anti-thrombotic mechanisms to maintain blood fluidity, including synthesis of a heparan sulfate rich glycocalyx that binds anti-thrombin (which inhibits thrombin), production of nitric oxide (which inhibits platelet activation), expression of tPA, and surface expression of TM (which diverts the pro-coagulant activities of thrombin to the anti-thrombotic functions of APC) (Figure 6A). These mechanisms are suppressed by inflammation, shifting the local environment towards a pro-thrombotic phenotype. Targeting ATA to the luminal surface of endothelium is intended to compensate for these pro-thrombotic changes without compromising normal tissue responses to injury.

Transfection of endothelium with genes encoding ATAs provided a prototype for such intervention. For example, transfected baboon endothelial cells (ECs) expressing tPA or a membrane-anchored form of uPA inhibited formation of platelet- and fibrin-rich thrombi in vitro, as did jugular vein segments transduced to express TM prior to graft insertion. However, the challenge of applying gene therapy to acute settings limits its utility in managing thrombosis.

Conjugating ATA to ligands that bind to the endothelial surface offers a potential alternative approach. Ideally, ATAs should anchor preferentially onto pro-thrombotic endothelium without cell injury or internalization of drug. For example, hirudin cross-linked to anti-E-selectin antibody binds to cytokine-activated ECs and inhibits thrombin in vitro but has limited therapeutic utility due to endocytosis in vivo. Likewise, anti-ACE/tPA conjugate is highly effective at targeting the endothelium in vivo but is endocytosed and has little fibrinolytic utility.
The endothelial adhesion molecules PECAM-1 and ICAM-1 are attractive targets for ATA delivery, respectively, to normal (i.e. prophylaxis) or activated (therapy) endothelium. Inhibition of their native functions may provide additional anti-inflammatory benefit. Although ECs internalize multivalent antibody conjugates targeting ICAM-1 and PECAM-1, they do not internalize monomolecular ligands. Accordingly, a series of ATA fused with monovalent scFv fragments of PECAM-1 antibodies have been devised and tested in animal models. Prophylactic injection of lmw-scuPA fused with anti-PECAM scFv enhanced lysis of subsequently formed emboli in the pulmonary vasculature. Similarly, prophylactic injection of this fusion via the carotid artery resulted in the rapid lysis of thromboemboli in the cerebral vasculature and attenuation of cerebral edema, whereas its untargeted ATA counterpart exacerbated brain injury.

PECAM-targeting of ATAs, which acts locally in a thrombin-dependent manner, has also been tested (Figure 6B). scFv/lmw-scuPA-T provided potent thromboprophylaxis against thrombin-induced pulmonary thrombosis, reduced pulmonary deposition of fibrin and improved oxygenation in a model of focal pulmonary ischemia/reperfusion injury. Endothelial targeting of scFv/TM may provide even greater opportunities for enhanced anti-coagulant and anti-inflammatory effects through interactions with the endothelial protein C receptor (EPCR). Ongoing preclinical studies are focused on segregating the diverse activities of scFv/TM, including quenching of cytokines and generation of thrombin-activatable thrombolysis inhibitor (TAFI), in addition to its anti-thrombotic activity.

4. Conclusion: challenges and opportunities. Emergency thrombolysis and thromboprophylaxis rely on the narrow margin between benefit and risk inherent in the use of existing ATAs. These drugs fail to discriminate between healthy and at-risk vasculature and distribute widely in the circulation. Recently developed ADDS, designed to profit from altered hemodynamic forces and target accessibility, might be capable of sensing subtle differences between mural hemostatic and occlusive thrombi. Such ADDS, including those using natural blood cells of the host as carriers, hold promise to enable Transient Targeted Thromboprophylaxis, a rapid and precise intervention capable of distinguishing pathological from hemostatic clots, and provide safe and effective management of acute thrombotic disorders not amenable to current treatment regimens (Figure 7).

By changing ATA delivery in a manner that concentrates effective doses in the vicinity of the target and provides a clinically appropriate temporal window, it might be possible to capitalize on advantages provided by the dynamic nature of clot initiation, propagation and perseverance. Most likely, clinical utility of targeted ATA requiring parenteral administration will be limited to acute settings, such as in the Emergency Department or Interventional Radiology suite.
Conditions that may be amenable to delivery of ATA (Supplemental Figure 4) include non-ST segment elevation-AMI, sub-massive pulmonary embolism, and acute mesenteric or limb ischemia. TTT might also find utility in postsurgical patients (Figure 1), as a means to prevent post-perfusion syndrome during cardiopulmonary bypass, and, perhaps, in patients suffering stuttering cerebral ischemia from recurrent in situ thrombosis or thromboembolism. Recombinant scFv/fusion proteins allow targeting of functional fragments of prodrugs to sites of ongoing thrombosis using diverse carriers and targets (RBC, endothelium, platelets, etc.) and activation options. Biological drugs such as recombinant fusions have been incorporated into clinical care and nanocarriers have been tested to treat oncological diseases.

Despite substantial preclinical data indicating a potential role for ADDS in the delivery of ATAs, enthusiasm must be tempered by several important factors. Targeted ATAs have not been tested in humans, and their translation to clinical use will be challenging. The animal models in which the majority of testing has occurred may not mirror important clinical features, especially the risk of bleeding. Clinical trial design is likely to be challenging due to the acute and unpredictable onset of the relevant thrombotic conditions. It might be difficult to find a comparison group of free drug in those settings in which we envision use of targeted ATAs and current thromboprophylaxis poses too high a risk. At the same time, any new agent will have to show superior efficacy, safety, or another compelling benefit over existing therapy, at a time in which multiple new small molecule ATAs are vying to replace older agents in routine clinical use. As with any new biotherapeutic, production costs may be higher than they are for synthetic ATAs and, at least initially, the regulatory landscape may be complex. However, if ATAs delivered via ADDS can demonstrate a meaningful benefit in patient outcome in well chosen clinical settings, then this approach is likely to overcome these hurdles and attract considerable interest from physicians and pharmaceutical companies alike.
Acknowledgments:

This work is supported by NIH grants HL087036, HL090697, and HL091950.

Authorship contributions: C.F.G., M.D.H., R.C., D.B.C. and V.R.M. wrote the manuscript.

Conflict-of-interest disclosure: The authors declare no competing financial interests.
REFERENCES

20. Wang S-S, Chou N-K, Chung T-W. The t-PA-encapsulated PLGA nanoparticles shelled with CS or CS-GRGD alter both permeation through and dissolving patterns of blood clots.


Table 1: Comparative analysis of the approaches for optimization of ATA pharmacokinetics. Each ADDS has distinct advantages and disadvantages. However, in general all suffer from impeded permeation into clots. Newer “smart” systems (SA-NTs, ELIPs) with mechanisms for achieving concentrated local drug delivery may have some success in overcoming this limitation. Alternatively, the use of “natural” delivery systems, such as RBCs, and their mimetics, may provide sufficiently prolonged circulation to allow for prophylactic use.

<table>
<thead>
<tr>
<th>Carrier</th>
<th>ATA Loading Method</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>~T₁/₂ in blood</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEG coating</td>
<td>Covalent chemical conjugation</td>
<td>Biocompatible; extensively studied in vitro and in animals; FDA-approved, clinically used for other drugs</td>
<td>Reduction of ATA activity; product heterogeneity</td>
<td>Minutes to hours</td>
</tr>
<tr>
<td>Liposome</td>
<td>Encapsulation</td>
<td>Same as PEG conjugates</td>
<td>Low encapsulation efficiency; limited stability and stealth features</td>
<td>Minutes to hours</td>
</tr>
<tr>
<td>Polymeric nanoparticles</td>
<td>Encapsulation or surface coating</td>
<td>Biodegradable; tunable properties (size, shape, release kinetics); stability</td>
<td>Harsh preparation procedures aggravate loss of activity; delayed drug release; potential toxicity; mostly in vitro studies</td>
<td>Minutes to hours</td>
</tr>
<tr>
<td>Dendrimers</td>
<td>Conjugation</td>
<td>Precise control of size and structure</td>
<td>Same as polymeric nanoparticles; low drug loading; laborious synthesis</td>
<td>Minutes to hours</td>
</tr>
<tr>
<td>Echogenic Liposome (ELIP)</td>
<td>Encapsulation</td>
<td>Biocompatible; imaging capacity; ultrasound-enhanced drug release</td>
<td>Same as liposomes, ultrasound-induced enzyme inactivation</td>
<td>Minutes to hours</td>
</tr>
<tr>
<td>Shear-activated nanotherapeutics (SA-NTs)</td>
<td>Conjugation</td>
<td>Biodegradable; shear-based local drug release in stenotic vessels</td>
<td>Early development stage; low conjugation efficiency; product heterogeneity; unknown whether fully occlusive clots can be attacked</td>
<td>NR</td>
</tr>
<tr>
<td>Filomicelles, RBC/platelet mimetics</td>
<td>ATA not yet loaded</td>
<td>May significantly prolong circulation of ATA</td>
<td>Untested for ATA delivery; limited data in animals; potential toxicity; complex production &amp; use</td>
<td>Hours to days</td>
</tr>
<tr>
<td>RBC</td>
<td>Conjugation</td>
<td>Significantly prolongs ATA circulation; no diffusion into tissues; no lysis of fresh preformed hemostatic clots; markedly reduced adverse effects of ATA in the CNS; efficacy and safety proven in diverse types of vascular thrombosis in several animal species</td>
<td>Need for ex vivo conjugation of ATA to isolated RBC and transfusion; cannot be used to dissolve existing occluding clots</td>
<td>Hours to days</td>
</tr>
</tbody>
</table>
Table 2. Approaches to vascular targeting of ATAs. The outline provides an historical perspective: from the oldest and most intensively studied approaches to target clot components to less well established but promising techniques to deliver ATAs to cellular targets in the vasculature. PECAM: platelet-endothelial cell adhesion molecule, ICAM: intercellular adhesion molecule, VCAM: vascular cell adhesion molecule, EC: endothelial cell, ACE: angiotensin converting enzyme.

<table>
<thead>
<tr>
<th>Target category</th>
<th>Targets</th>
<th>Goal/Advantages</th>
<th>Challenges</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clot components</td>
<td>Fibrin, Coagulation factors, Activated platelets</td>
<td>Concentrate ATA in occlusive or growing clots; inhibit pro-thrombotic functions of targets</td>
<td>Targets rapidly disappear or become inaccessible in clots; clot permeation; short PK restricts prophylactic utility</td>
</tr>
<tr>
<td>Blood cells</td>
<td>Resting platelets</td>
<td>Use natural tropism for the above goal; inhibit pro-thrombotic functions of platelets</td>
<td>Limited animal data on PK, targeting and effects; restricted access to and effect on preformed occluding clots</td>
</tr>
<tr>
<td></td>
<td>RBC</td>
<td>Significantly prolonged PK and restricted side effects of RBC-bound ATAs enable safe TTT in animal models</td>
<td>Restricted access to and effect on preformed occlusive clots; animal studies are limited to mouse models</td>
</tr>
<tr>
<td>Endothelium</td>
<td>PECAM</td>
<td>Prophylactic anchoring of ATAs on EC preconditions to subsequent thrombosis</td>
<td>No/poor access for EC-targeted ATA to EC in sites of preformed/ongoing thrombosis</td>
</tr>
<tr>
<td></td>
<td>ICAM</td>
<td>As above; enriched anchoring of ATAs in inflamed EC; inhibition of WBC traffic</td>
<td>As above</td>
</tr>
<tr>
<td></td>
<td>Selectins and VCAM</td>
<td>As above; high selectivity to inflamed EC</td>
<td>As above; transient target exposure and fast internalization limit amplitude and duration of action</td>
</tr>
<tr>
<td></td>
<td>ACE</td>
<td>Enriched anchoring in the pulmonary vasculature</td>
<td>As in PECAM; fast internalization</td>
</tr>
</tbody>
</table>
FIGURE LEGENDS

Figure 1. Transient thromboprophylaxis (TTT) for prevention of post-surgical thrombosis. Thromboprophylaxis is especially problematic in the early post-operative period. Thrombosis is a common complication of the trauma, inflammation and immobility that accompany surgery, and thrombi are prone to recur or extend within hours of fibrinolysis, necessitating protracted activity. However, the risk of bleeding may preclude intervention for hours to days. Fibrinolytics are not used for prophylaxis because they are rapidly inactivated and cleared from the blood, necessitating use of high and unsafe doses to maintain activity. A rapid transient intervention that spares hemostatic clots, but prevents subsequent development of occlusive thrombosis elsewhere would fill a clear void in clinical management. This is but one of many settings in which patients at known imminent risk of thrombosis would benefit from TTT. An ideal agent for TTT would have an immediate onset of action that is of sufficient duration to prevent occlusive thrombi from forming without affecting pre-existing fresh hemostatic clots (e.g., those formed in the wound within ~20-30 min of uncomplicated surgery) and without causing off-target toxicity.

Figure 2. Advanced drug delivery systems (ADDS) for ATA. The schematics present examples of classes of carriers tested for delivery of ATA, their principal structure, and PA cargoes. Synthetic carriers vary in size from few nanometers to several microns, shape, stability and principle of drug loading and release. Liposomes and PEG represent the two most intensively studied ADDS for various ATAs including anticoagulants, platelet antagonists and listed fibrinolytics.

Figure 3: Coupling plasminogen activators to carrier RBC may convert risky fibrinolytic therapy into safe transient thromboprophylaxis. Upper panel: plasminogen activators (PAs) are used in the emergency setting for reperfusion. High doses are needed to compensate for rapid inactivation by PAI-1 and clearance by the liver and other organs. PAs diffuse into hemostatic mural clots as well as occlusive pathological clots, increasing the risk of bleeding, and into tissues, such as the CNS, where they cause toxicity. Therapeutic doses can deplete fibrinogen and plasminogen increasing the risk of bleeding while reducing fibrinolytic potential. Only a small proportion of the residual circulating PA reaches and penetrates the interior of clots. Lower panel: RBC-coupled PAs avoid rapid clearance, circulating for many hours to days without damaging carrier RBCs. The large size and hemodynamic factors prevent access to pre-existing hemostatic clots, limits extravasation into tissues and blocks interaction with cellular receptors such as uPAR (CD87) and integrins that may initiate deleterious intravascular signaling cascades. RBC-coupled PAs are incorporated within nascent intravascular clots formed after treatment, which they lyse from within, rapidly restoring flow and preventing ischemia at doses that are orders of magnitude less than free fibrinolytics.

Figure 4. Strategies for ATA targeting. The action of clot-targeted scFv/ATA has been studied in animal models of arterial and venous thrombosis. ATA fusions or antibody conjugates injected in the vasculature shortly after thrombotic occlusion (T3) circulate for a limited period of time. Affinity to clot components enhances the fraction of injected dose ATA accumulating at the thrombotic site. Fibrin-targeted ATAs bind to fibrin accessible at the clot surface and to additional fibrin depositing in the site of thrombosis in the course of clot growth and remodeling (light gray area surrounded by dash line). With similar interventions, ATAs bind to activated platelets accessible from clot surface and in remodeling clot, or to resting platelets (which may prolong circulation and provide continuous platelet-mediated delivery into secondary or remodeling clots). Clot-associated anticoagulants (e.g., hirudin) inhibit additional formation of thrombin, thus limiting thrombus growth, whereas plasminogen activators facilitate dissolution of clots and foster reperfusion.

Figure 5. Transient Targeted Thromboprophylaxis by RBC-targeted fusion ATA pro-drugs. ATAs, including fibrinolytics such as plasminogen activators (tPA and uPA) and anticoagulants such as thrombomodulin (TM) fused with an anti-RBC scFv bind to RBC in bloodstream circulate for a prolonged time without damaging the cell carrier or causing other adverse effects typical of soluble drugs (illustrated in Figure 5). Using RBC-targeted fusion constructs avoids the need for ex vivo loading and transfusion, thereby increasing clinical utility. Altering the mode of enzymatic activation from plasmin to thrombin enhances safety and localization. Upper panel: scFv/uPA-T containing a variant of uPA sensitive to
activation by thrombin exerts maximal fibrinolytic activity at sites of active clotting where the concentration of thrombin is greatest. Lower panel: scFv/TM facilitates local conversion by thrombin of protein C into APC, which in turn inactivates activated coagulation factors Va and VIIIa and inhibits pro-thrombotic and pro-inflammatory signaling via protease activated receptor (PARs) on platelets and endothelial cells, respectively.

**Figure 6. Endothelial Targeting of ATA.** Upper panel: Healthy endothelial cells (ECs) express antithrombotic activity, utilizing a variety of anti-platelet, anticoagulant, and fibrinolytic mechanisms to maintain blood fluidity. Lower panel: Thrombin-dependent therapeutics, including lmw-scuPA-T and TM, can be anchored to the luminal membrane of ECs using PECAM-targeted scFv antibody fragments. This approach has the potential to augment endogenous antithrombotic mechanisms, which developed through millions of years of evolution in order to prevent thrombosis without compromising normal tissue responses to injury.

**Figure 7. Advanced drug delivery systems cover a new area of anti-thrombotic utility: Transient Targeted Thromboprophylaxis.** Using ADDS has the potential to provide ATA pro-drugs with an immediate onset (local activation) and a duration of action lasting hours to days, targeted carriers that capitalize on the dynamic nature of thrombi and differences between preformed hemostatic and subsequently formed occlusive clots. This approach might extend the medical utility of fibrinolytic or anticoagulant ATAs.
FIGURES

Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6

Diagram showing the interaction between different molecules and pathways. The diagram includes labels for glycocalyx, ATIII, NO, eNOS, tPA, Pn, Pg, TM, PC, APC, PECAM-1, Anti-PECAM scFV/uPA-T, and Anti-PECAM scFV/TM.
Figure 7
Advanced drug delivery systems for anti-thrombotic agents

Colin Greineder, Melissa D. Howard, Ron Carnemolla, Douglas B. Cines and Vladimir Muzykantov