THROMBOSIS AND HEMOSTASIS

Targeted mutagenesis of zebrafish antithrombin III triggers disseminated intravascular coagulation and thrombosis, revealing insight into function

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Key Points

- Juvenile zebrafish tolerate widespread coagulopathy due to complete ablation of antithrombin III, but develop lethal thrombosis as adults.
- In vivo structure/function analysis of antithrombin III in zebrafish reveals limited roles for heparin-binding and anti-IXa/Xa activity.

Pathologic blood clotting is a leading cause of morbidity and mortality in the developed world, underlying deep vein thrombosis, myocardial infarction, and stroke. Genetic predisposition to thrombosis is still poorly understood, and we hypothesize that there are many additional risk alleles and modifying factors remaining to be discovered. Mammalian models have contributed to our understanding of thrombosis, but are low throughput and costly. We have turned to the zebrafish, a tool for high-throughput genetic analysis. Using zinc finger nucleases, we show that disruption of the zebrafish antithrombin III (at3) locus results in spontaneous venous thrombosis in larvae. Although homozygous mutants survive into early adulthood, they eventually succumb to massive intracardiac thrombosis. Characterization of null fish revealed disseminated intravascular coagulation in larvae secondary to unopposed thrombin activity and fibrinogen consumption, which could be rescued by both human and zebrafish at3 complementary DNAs. Mutation of the human AT3-reactive center loop abolished the ability to rescue, but the heparin-binding site was dispensable. These results demonstrate overall conservation of AT3 function in zebrafish, but reveal developmental variances in the ability to tolerate excessive clot formation. The accessibility of early zebrafish development will provide unique methods for dissection of the underlying mechanisms of thrombosis. (Blood. 2014;124(1):142-150)

Introduction

Venous thrombosis affects approximately 1 in 1000 individuals, resulting in over 300,000 hospitalizations per year. There are many well-described coagulation factor defects which are associated with thrombosis, including deficiencies of the natural anticoagulant proteins, protein C (PROC), protein S (PROS1), and antithrombin III (AT3).1 Homozygous deficiency of PROC or PROS1 is exceedingly rare and results in neonatal purpura fulminans in humans.1 However, mouse knockout models display more severe phenotypes, with neonatal and embryonic loss for PROC and PROS1 deficiencies, respectively.2-4 Complete loss of AT3 has not been described in humans, and is presumed to be in utero lethal.1 This is supported by the observation that homozygous deletion of AT3 in mice results in prenatal lethality, with widespread fibrin deposition in the heart and liver.5 AT3-deficient embryos also exhibit paradoxical hemorrhage, suggesting a consumptive coagulopathy,5 but this has been difficult to study given the relative inaccessibility of mammalian embryogenesis.

AT3 (also known as SERPINC1) is a serpin (serine protease inhibitor) that regulates many coagulation proteases including thrombin and factors IXa and Xa,6,7 and consists of 2 functional domains. The reactive center loop (RCL) presents the Arg393-Ser394 bond for cleavage, resulting in the formation of a stable inactive AT3-protease complex. The heparin-binding site binds endogenous glycosaminoglycans or infused heparin, acting as a bridge to the protease and inducing allosteric changes in AT3, which accelerates inhibition.6,7 Deficiency of AT3 is classified into 2 categories: type I with reduced plasma antigen levels and type II with functional defects, but normal AT3 levels.1 The latter is further subdivided into reactive site, heparin binding, and pleiotropic defects. Over 200 different mutations from patients with AT3 deficiency have been identified,8 and rapid advances in sequencing technologies guarantee that there are many more to come. Although in vitro assays are available for assessment of AT3 function, in some patients the underlying mechanism of AT3 deficiency remains unknown.

Teleost fish possess highly conserved orthologs of nearly all blood coagulation factors.9-12 Zebrafish (Danio rerio) have been widely used to study hemostasis, demonstrating conservation and function of the structural components of coagulation, including


The online version of this article contains a data supplement.

There is an Inside Blood Commentary on this article in this issue.
Figure 1. Targeted disruption of at3 using genome editing nucleases results in adult lethality. (A) ZFN-induced deletions in somatic cells. Of 16 pooled morphologically normal embryos from 2 individual experiments, 3 of 192 sequences were identified with ΔS (2x) and ΔG (1x) in exon 5. The top line is the at3 reference genome sequence from an unrelated individual. The blue and green arrows indicate 5’ and 3’ ZFN-binding sites, respectively. Deletions are indicated by red dashes. (B) Sequences of ZFN-induced mutations transmitted through the germline. Red dashes and letters indicate deletions and insertions, respectively. (C) Survival curve of a total of 3 clutches derived from at3+/− heterozygous crosses shows loss of homozygotes. Offspring were genotyped at 2 to 4 months of age and tracked over 1 year. The results demonstrate a highly significant difference in the survival of homozygous mutants by log-rank (Mantel-Cox) analysis (P < .0001). (D) Externally visible secondary hemorrhage in an at3 homozgyous mutant is indicated by an arrow. This fish was identified while alive, anesthetized, and the image was captured using a Leica MZ16FA microscope and a Canon EOS6D camera. (E-F) RT-PCR analysis of at3 variants demonstrates reduction of at3 expression in heterozygotes and absence of homozygous mutant mRNA. (E) Top: Total mRNA was prepared from adult fish (4 mpf) followed by qualitative RT-PCR. Bottom: Genotyping of at3 mutants. (F) Total mRNA was prepared from pooled larvae (3 dpf; n = 15 for each genotype) followed by quantitative real-time PCR. Error bars represent standard deviation and statistical significance was determined by t test. (G) at3 homozygous mutants were injected with a ubi regulated at3 cDNA expression vector and followed for 1 year.

Methods

Animals

Zebrafish were raised in accordance with animal care guidelines as approved by the University of Michigan Animal Care and Use Committee. Embryos are defined as 0 to 2 days postfertilization (dpf), and larvae as 3 to 29 dpf.

Targeted mutagenesis using ZFNs

ZFN pairs were designed to target exon 5 of the at3 genomic locus. ZF arrays were used to construct ZFNs (Oligomerized Pool ENgineering) as described.22,23 Targeted mutagenesis with the constructed ZFNs was completed essentially as described.19,24 ZF arrays were cloned into expression vectors pMLM290 and 292, and transcribed in vitro using T7 mMESSAGE mMACHINE (Ambion). One nanogram of each left and right ZFN messenger RNA (mRNA) was injected into 1-cell AB X TL F1 embryos, as described.14 At 2 to 3 dpf, 8 morphologically normal embryos were pooled and lysed in buffer (10 mM Tris, pH 8.0, 200 mM NaCl, 10 mM EDTA, 0.5% sodium deoxycholate [SDS], 100 μg/mL Proteinase K) at 50°C for 2 to 3 hours, and genomic DNA isolated by phenol:chloroform extraction and ethanol precipitation. Polymerase chain reaction (PCR) of the target site was performed, and products cloned into pCR-Blunt (Invitrogen). PCR of 96 colonies was performed using M13 reverse and T7 primers, and products sequenced. The remaining injected F0 embryos were raised to adulthood and crossed to wild-type fish. Sixteen to 24 F1 embryos from each F0 founder were screened for ZFN-induced mutations by PCR of the target site with a 6-FAM–labeled gene-specific primer, after lysis in PCR extraction buffer24 (10 mM Tris pH 8.0, 2 mM EDTA, 0.2% Triton X-100 and 100 μg/mL Proteinase K). PCR products were resolved by capillary electrophoresis on a 3730XL DNA Analyzer (Applied Biosystems). Data were processed using GeneMarker (Softgenetics LLC) for mutant identification. The remaining F1 offspring were raised to adulthood and heterozygotes identified by fin clip genotyping.

Genotyping of mutant offspring

Fin clip biopsies were obtained from adult fish after anesthesia in tricaine (0.16 mg/mL; Western Chemical Inc), and larvae were humanely killed in high-dose tricaine (1.6 mg/mL), followed by lysis in PCR extraction buffer.24 Deletion mutations were detected by PCR and agarose gel electrophoresis (Figure 1E bottom). Primers were designed using Primer325 and oligonucleotides ordered from Integrated DNA Technologies. All oligonucleotide sequences are listed in supplemental Tables 1-2 (see supplemental Data available on the Blood Web site).

Construction of at3 in vivo expression vectors

Vectors pubi-zat3-EGFP and pubi-hAT3-EGFP were constructed by inserting complementary DNA (cDNA) sequences under control of the ubi promoter in
frame with enhanced green fluorescent protein (EGFP) in pDestTol2pA2_ubi EGFP29 (supplemental Figure 1). Zebrafish at3 was amplified from total adult cDNA using NcoI tagged primers, and cloned into NcoI digested pDestTol2pA2_ubi:EGFP. Human AT3 was amplified from liver cDNA using primers designed for sequence- and ligation-independent cloning, and cloned into NcoI digested pDestTol2pA2_ubi:EGFP as described.27

Zebrafish and human at3 substitutions were introduced into pubi-at3-EGFP and pubi-hAT3-EGFP, respectively, as described with some modifications.28 Mutagenic primers amplified the plasmids using Phusion DNA polymerase (New England Biolabs) as follows: 98°C for 3 minutes; 4 cycles of 98°C for 10 seconds, 55°C for 30 seconds, 72°C for 8 minutes; 12 cycles of 98°C for 10 seconds, 60°C for 30 seconds, 72°C for 8 minutes; 10 cycles of 98°C for 10 seconds, 55°C for 30 seconds, 72°C for 10 minutes. PCR products were column purified, digested with DpnI, and transformed into TOP10 cells, and mutations confirmed by sequencing. Amino acid numbering of human AT3 begins at +1 after the propeptide cleavage site (amino acid 33 from the initiation methionine), reflecting absence of the 32-aa propeptide. The N-terminal amino acids of zebrafish At3 are not conserved with the mammalian AT3 propeptide (supplemental Figure 2), therefore, +1 for the former is from the initiation methionine.

Thrombin injection
Four milliunits of bovine thrombin (Sigma-Aldrich) were retro-orbitally infused33 into 4 dpf larvae from at3Δ90/Δ90 intercrosses. Phenotypes were collected 2 minutes postinjection by observers blinded to genotype. After phenotyping, larvae were lysed and genotyped (see supplemental Methods for a detailed description).

Laser-mediated endothelial injury
Endothelial injury was executed using a pulsed nitrogen dye laser and focusing system (MicroPoint; Andor Technology) as described.14,17 Larvae (3-4 dpf) were anesthetized in tricaine, embedded in 0.8% low-melt agarose on glass cover slips, and visualized on an inverted microscope (Olympus IX71, 40× objective). The posterior cardinal vein (PCV) endothelium was targeted and laser-ablated at the fifth somite distal from the anal pore at power level 18 using 100 pulses. The time to occlusion was recorded up to 2 minutes. Larvae were recovered from agarose and genotyped.

Fibrinogen injection and ε-aminocaproic acid treatment
Offspring from at3Δ90/Δ90 and at3Δ90/Δ90 crosses were either infused with 50 ng (2 nL) human fibrinogen or bovine serum albumin (Sigma-Aldrich) in saline at 3 dpf, or preincubated with 0.1M ε-aminocaproic acid at 2 dpf, and laser injury performed at 60 minutes postinjection or 24 hours postincubation, respectively, followed by genotyping.

FITC-fibrinogen labeling and infusion
Human fibrinogen was labeled with fluorescein isothiocyanate (FITC; Thermo-Scientific), PD-10 column purified, and dye-to-protein molar ratio determined by manufacturer’s instructions. Eight to 20 ng of a 1:1 molar ratio of FITC:fibrinogen was infused at 3 dpf.

Embryos were incubated in 50 μg/mL warfarin (Sigma-Aldrich) for 48 hours prior to FITC-fibrinogen infusion, or larvae infused with 1 mg/mL tissue plasminogen activator (tPA; Alteplase) via retro-orbital injection. At 3 dpf, larvae were infused with FITC-fibrinogen and evaluated 1 hour postinjection or postinfusion with tPA, and images were acquired with a charge-coupled device camera. Fluorescence intensity was determined either qualitatively by a blinded observer or quantitatively using Scion/ImageJ imaging software.34 Corrected total cell fluorescence was integrated density after subtraction of background (area selected × mean background fluorescence).35

RNA isolation, generation of cDNA, and reverse transcription PCR
Larvae (3 dpf) were anesthetized in tricaine and biopsied as described16 for genotyping. Larvae were maintained in RNAlater (Ambion) and 15 of each genotype pooled for RNA extraction. Total RNA was isolated using the PureLink RNA isolation kit (Ambion), and reverse transcribed with Superscript Synthesis (Invitrogen). cDNAs were amplified (MyQ; Bio-Rad) using SYBR Green Master Mix (Applied Biosystems) at 95°C for 10 minutes, 40 cycles of 95°C for 30 seconds, 55°C for 30 seconds. Each was analyzed in triplicate and normalized to 18S ribosomal RNA, and expression presented as fold changes (ΔΔCt cycle threshold (ΔΔCt)).36

Histologic examination
Humanely killed adult zebrafish were fixed in 4% buffered formalin overnight at room temperature, bisected in the midsagittal plane, embedded in paraffin, and sagittal sections produced at least 50 μm apart. Sections (3-4 μm) were hematoxylin and eosin stained.

Statistical analysis
Analyses were performed using χ², Mann-Whitney U, Fisher exact, or 2-tailed Student t tests. Log-rank testing of survival curves was performed using Prism (GraphPad Software). For statistical comparisons between individual transgenic rescue experiments, results from homozygous mutants were normalized to the median of their heterozygous clutchmates. P values
were calculated with the Mann-Whitney U test by comparing the normalized data, followed by Bonferroni correction.

**Results**

**Targeted disruption of at3 using genome editing nucleases results in adult lethality in association with massive intracardiac thrombosis**

We used OPEN\textsuperscript{19,22,23} to produce 4 ZFN pairs designed to cleave exon 5 of the zebrafish at3 genomic locus (zebrafish nomenclature guidelines dictate use of “at3” for the gene, “At3” for protein\textsuperscript{38}). RNA encoding each ZFN pair was injected into the cytoplasm of single-cell embryos, with 1 pair producing somatic mutations (Figure 1A) at a frequency of 1.6% (3 of 192 sequenced clones from 2 pools of 8 embryos). The remaining embryos were raised to adulthood, and 46 of these F0 at3\textsuperscript{+/−} were mated to wild-type fish to confirm germline transmission. We identified 4 founders with the following mutations: 4-bp insertion (at3\textsuperscript{+/+}), 6-bp deletion (at3\textsuperscript{−/−}), 22-bp deletion (at3\textsuperscript{Δ22}), and 90-bp deletion (at3\textsuperscript{Δ90}) (Figure 1B). at3\textsuperscript{−/−}, at3\textsuperscript{Δ22}, and at3\textsuperscript{Δ90} produced frameshifts in exon 5 and are predicted to result in null alleles (the 90-bp deletion removes the splice donor site) (supplemental Table 3). The expected Mendelian distribution (Table 1) with no visible phenotypes. However, beginning at ~2 months postfertilization (mpf), we observed a significant loss of homozygotes (Table 1), with ~80% dying between 2 and 7 mfp (Figure 1C). Approximately 5% of at3\textsuperscript{−/−} mutants had externally visible hemorrhage (Figure 1D). Reverse transcription PCR (RT-PCR) indicated reduction of at3 expression in heterozygotes, and absence of expression in homozygous mutants, presumably due to nonsense-mediated decay\textsuperscript{39} (Figure 1E-F).

To rule out potential off-target ZFN mutations, we injected at3\textsuperscript{Δ22} incrosses with an at3 cDNA under control of the zebrafish ubiquitin (ubi) promoter (ubi-at3\textsuperscript{Δ22}). After 1 year, homozygotes displayed significantly greater survival compared with uninjected at3\textsuperscript{−/−} fish (Figure 1G).

Histologic examination of surviving at3\textsuperscript{−/−} fish at 2 to 3 mfp revealed large intracardiac thrombi in all homozygous mutants examined (n = 10), but not in wild-type (n = 10) or heterozygous mutants (n = 10) (Figure 2A-D). Notably, evaluation of a ubi-at3 rescued homozygous at3\textsuperscript{Δ22} mutant found no evidence of thrombosis at 16 mfp (supplemental Figure 3).

Table 1. Genotyping results of heterozygous incrosses from mutant line at3\textsuperscript{Δ90}

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at3 homozygous mutants display an increased time to occlusion in an induced model of venous thrombosis

Complete deficiency of AT3 in mice and humans results in lethal thrombosis in utero.\textsuperscript{1,5} Although loss of zebrafish At3 resulted in apparently lethal thrombosis in adults, embryos and larvae appeared grossly normal with no overt evidence of pathologic thrombosis. We performed laser injury\textsuperscript{14,17} at 3 to 4 dpf on larvae derived from at3\textsuperscript{+/−} intercrosses, targeting the posterior cardinal vein (PCV) (orthologous to the mammalian inferior vena cava; supplemental Figure 4 and supplemental Video). As expected, wild-type and at3\textsuperscript{+/−} larvae exhibited vessel occlusion within 2 minutes, but at3\textsuperscript{−/−} mutants failed to occlude within that time period (Figure 3A). To confirm the role of At3 deficiency in this bleeding phenotype, we injected ubi regulated human and zebrafish at3 cDNAs into single-cell embryos of wild-type and at3\textsuperscript{−/−} clutches, followed by laser injury at 3 dpf. Fifty-two and 83% of human and zebrafish transgene-injected embryos, respectively, demonstrated a time to occlusion of <2 minutes (P < .0001 by the Mann-Whitney U test, Figure 3B-C).

The unexpected prolongation of clotting time in at3\textsuperscript{−/−} larvae suggested the possibility of differential localization of zebrafish At3 compared with its mammalian ortholog. However, in situ hybridization at 5 dpf revealed strong mRNA expression localized predominantly to the liver, consistent with mouse and human AT3\textsuperscript{30,41} (Figure 4A-D). Minor expression of At3 has been noted in the notochord and neural tube of embryonic day 10.5 mice,\textsuperscript{42} whereas early zebrafish embryos demonstrate weak yolk syncytial layer expression from 16 through 48 hours postfertilization.\textsuperscript{43} Alignment with human AT3 revealed that the RCL was highly conserved across species (supplemental Figure 2). The orthologous P1 arginine was identified (Figure 4E), substituted with glycine (zebrafish Arg410Gly, orthologous to human Arg393), and assessed for binding to human thrombin in vitro (Figure 4F). Although wild-type At3 exhibited binding to thrombin and formed a putative thrombin-antithrombin complex, this interaction was absent in reactions with At3\textsuperscript{R410G} (Figure 4F). To prove that the mutant phenotype was dependent on known serine protease activity, an At3\textsuperscript{R410G} expressing cDNA was injected into at3\textsuperscript{+/−} embryos. Consistent with the in vitro–binding studies, At3\textsuperscript{R410G} failed to rescue the at3\textsuperscript{−/−} larval bleeding phenotype (Figure 4G). Taken together, these data strongly suggest that the observed at3\textsuperscript{−/−} larval phenotype is due to loss of anticoagulant activity.
Hypofibrinogenemia in at3−/− mutants is secondary to DIC

We hypothesized that the mechanism underlying the discrepant adult thrombotic and larval bleeding phenotypes in at3 mutants was DIC, with microscopic fibrin deposition. To identify the location of purported fibrin, human fibrinogen was labeled with FITC44 and infused into at3+/− intercross progeny. Widespread PCV fluorescence was observed in at3 homozygous mutant larvae after infusion, but absent in wild-type and heterozygous siblings (Figure 6A-E). Warfarin prevented laser-induced thrombosis in wild-type larvae (supplemental Figure 5), and pretreatment of at3−/− larvae eliminated fluorescence accumulation (Figure 6F). Coinjection of tPA partially prevented accumulation, as did postinjection (Figure 6G-H). Taken together, these data support the presence of DIC with consumption of fibrinogen in at3−/− mutants.

Evaluation of human AT3 mutations in vivo

We examined several additional well-described human AT3 mutations associated with thrombosis, R393H,45-49 A404D,8 L99F,50,51 K114E,52,53, as well as a variant with greatly reduced anti-IXa and anti-Xa activity, Y253A4 (Figure 7). The P1 arginine mutant R393H was unable to reverse the DIC in at3−/− larvae. Surprisingly, the RCL mutant A404D and heparin-binding mutants L99F and K114E all demonstrated significant rescue, although the former was less potent than the latter two. Human AT3 Y253A also reversed the bleeding phenotype and was significantly more efficient when compared with human AT3 (P < .001).

Discussion

We report targeted mutagenesis of zebrafish at3 via ZFNs, one class of genome editing nuclease. Our data demonstrate that zebrafish survive complete loss of At3 with rampant DIC in the embryonic and larval periods, but apparently succumb to massive intracardiac thrombosis in adulthood. Concordant with our data, embryonic At3−/− mutant mice display a consumptive coagulopathy with secondary hemorrhage, although this causes in utero thrombosis and subsequent demise.2 It is notable that in both the complete mouse knockout and a heparin-binding mutant knockin (Arg48Cys),55 the heart is one of the most prominent sites of spontaneous thrombosis. The structure of the adult zebrafish heart has been described in detail,56 and overall cardiovascular development is highly comparable to mammalian and avian species. Taken together, our data support the contention that although the timing of ultimate lethal thrombosis is different between zebrafish and mammals, the underlying mechanisms of the coagulation cascade are conserved.

Our studies indicate that loss of At3 leads to dysregulation of coagulation, resulting in consumption of fibrinogen and prolonged...
clotting times in at3 homozygous mutant larvae. Although there were clearly visible spontaneous macrovascular thrombi in adult fish, injection of fluorescent fibrinogen identified disseminated microvascular thrombi in larvae, the latter consistent with clinical DIC. In patients, DIC is characterized by unregulated and excessive thrombin generation, resulting in fibrin formation with secondary plasmin activation and fibrinolysis. Clinical bleeding and/or thrombosis develops depending on the balance between thrombin and plasmin generation. In at3−/− larvae, α-aminocaproic acid, an inhibitor of plasmin, rescued the bleeding phenotype, demonstrating conservation of the fibrinolytic system. This also signifies possible contributions of fibrinolysis in this model, consistent with human DIC. These data demonstrate that loss of At3 results in unopposed procoagulant activity and possible secondary fibrinolytic activity, confirming a conserved role for zebrafish At3 as a key regulator of blood coagulation. Notably, extensive intracranial and subcutaneous hemorrhage was observed in

Figure 5. Evidence for fibrinogen consumption in at3 homozygous mutant larvae and adults. All data shown are from 190 mutants. (A) Thrombin injection of larvae derived from at3+/− increases at 4 dpf. Bar graph represents percentage of each thrombosis grade. Grade 0, normal flow within the PCV and dorsal aorta; grade 1, partial obstruction of flow in the PCV and dorsal aorta due to thrombus formation; grade 2, complete obstruction of flow. Statistical significance was determined by Fisher exact test. (B) Top: Western blot analysis of fibrinogen in adult plasma isolated from at3+/+ and at3−/− clutchmates from 3 separate experiments, n = 3 per genotype. Error bars represent standard deviation and statistical significance was determined by a paired t test. (C) Quantitative real-time PCR demonstrates no significant difference of fga and fgb mRNA expression in at3+/+ and at3−/− clutchmates (3 dpf, n = 15 per genotype). Error bars represent standard deviation and statistical significance was determined by a paired t test. (D) Rescue of time to laser-induced occlusion by human fibrinogen injection at 3 dpf. at3−/− mutant clutchmates were injected with human fibrinogen or BSA. Uninjected (n = 13), BSA injection (n = 9), and human fibrinogen injection (n = 33). (E) Effects of EACA (100 mM) pretreatment on at3−/− larvae. Untreated (n = 34), EACA treated (n = 36). Circles represent individual larvae. Statistical significance was determined by the Mann-Whitney U test. Horizontal bars represent the median time to occlusion. BSA, bovine serum albumin; EACA, α-aminocaproic acid; ns, nonsignificant.
mouse At3−/− embryos, without detectable accumulation of fibrinogen in those tissues.5 The authors posited that a consumptive coagulopathy depleted fibrinogen from the plasma, resulting in the observed bleeding, as we discerned in at3−/− mutants. Similarly, Proc knockout mice displayed intracranial hemorrhage in addition to thrombi in the heart and other organs, and clottable fibrinogen was undetectable.2 Recently, fga was targeted in zebrafish using ZFNs, and the authors demonstrated complete absence of fibrinogen.58 They did not observe specific bleeding in homozygous mutant larvae, but did note sporadic adult hemorrhage, both consistent with our data (Figure 1D). In contrast, they only detected a partial reduction in survival of homozygous mutants, supporting our data that suggest at3−/− mutants die of thrombosis rather than hemorrhage.

The observation that spontaneous, nonocclusive thrombosis due to DIC is so well tolerated in fish embryos and larvae raises important questions about the role of the coagulation system in vertebrate development. It is tempting to speculate that additional biological mechanisms may play a role, either through established clotting or other pathways. There may be protective species-specific factors for critical determinants of hemostatic balance, and the ability to perform high-throughput genetic screens in the zebrafish could lead to the identification of novel modifying factors.18 Another powerful application for zebrafish is small-molecule screening, which has led to the rapid clinical development of a novel therapeutic approach for cord blood transplant.59-61 Screens for novel anticoagulants or modulators of DIC might also have the potential to lead to innovative therapeutic modalities.

Alignment of human to zebrafish At3 reveals 55% identity and 70% similarity (supplemental Figure 2 and Kumar et al62). Because human AT3 rescued the prolonged time to occlusion in our endothelial injury assay, we performed semiquantitative evaluation of various human mutations and targeted substitutions. Substitution of the RCL P1 arginine (Arg393His) has been identiﬁed in several individuals and families with thrombophilia,45-49 and has been shown to alter AT3 function in vitro.63 Arg393His abolished AT3 activity, confirming the results of biochemical analysis as well as demonstrating the ability of this in vivo model to evaluate human coagulation factor variants. Ala404Asp, a substitution in the RCL coagulation factor variants. Ala404Asp, a substitution in the RCL
IXa- and Xa-binding exosite.\textsuperscript{54} Surprisingly, AT3\textsuperscript{Tyr253Ala} rescued the \textit{at3}\textsuperscript{-/-} coagulopathy with a greater potency than wild type. Although only semiquantitative, these data suggest that proper hemostatic balance primarily depends on thrombin inhibition. Also, AT3\textsuperscript{Tyr253Ala} exhibits enhanced rates of thrombin inhibition in the presence of heparin,\textsuperscript{54} which could explain the increased activity. Taken together, these data provide novel insights into AT3 function in vivo. Our approach establishes a rapid and robust new paradigm for validating human genetic variants associated with coagulation disorders, as well as complements biochemical and structure/function studies.

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References


J.A.S. is the Diane and Larry Johnson Family Scholar of Pediatrics and Communicable Diseases.

Authorship

Contribution: Y.L. designed and performed research, analyzed data, and wrote the manuscript; C.A.K. designed and performed research, and analyzed data; M.L.M. and J.K.J. designed and performed research; C.E.R., P.T., M.C.H., A.H.V., T.R., and Z.H. performed research; R.M. analyzed data; S.T.O. designed research and analyzed data; and J.A.S. designed, performed, and supervised research, analyzed data, and wrote the manuscript.

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