A sensitized mutagenesis screen in Factor V Leiden mice identifies novel thrombosis suppressor loci

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Abstract

Factor V Leiden $(F5^L)$ is a common genetic risk factor for venous thromboembolism in humans. We conducted a sensitized ENU mutagenesis screen for dominant thrombosuppressor genes based on perinatal lethal thrombosis in mice homozygous for $F5^L$ ($F5^{L/L}$) and haploinsufficient for tissue factor pathway inhibitor (Tfpi^{+/-}). F8 deficiency enhanced survival of F5^{L/L} Tfpi^{+/-} mice, demonstrating that $F5^{L/L}$ $Tfpi^{+/r}$ lethality is genetically suppressible. ENU-mutagenized $F5^{L/L}$ males and $F5^{L/L}$ females were crossed to generate 6,729 progeny, with 98 $F5^{L/L}$ Tfpi^{+/-} offspring surviving until weaning and 16 lines exhibiting transmission of a putative thrombosuppressor to subsequent generations. These lines are referred to as MF5L (Modifier of Factor 5 Leiden) 1-16. Linkage analysis in MF5L6 identified a chromosome 3 locus containing the tissue factor gene (F3). Though no ENU-induced F3 mutation was identified, haploinsufficiency for F3 ($F3^{+/-}$) suppressed $F5^{L/L}$ $Tfpi^{+/-}$ lethality. Whole exome sequencing in MF5L12 identified an Actr2 gene point mutation (p.R258G) as the sole candidate. Inheritance of this variant is associated with suppression of $F5^{L/L}$ $Tfpi^{+/-}$ lethality (p=1.7x10⁻⁶), suggesting that Actr2^{p.R258G} is thrombosuppressive. CRISPR/Cas9 experiments to generate an independent Actr2 knockin/knockout demonstrated that Actr2 haploinsufficiency is lethal, supporting a hypomorphic or gain of function mechanism of action for Actr2^{p,R258G}. Our findings identify F8 and the Tfpi/F3 axis as key regulators in determining thrombosis balance in the setting of F5^L and also suggest a novel role for Actr2 in this process.

Introduction

Venous thromboembolism (VTE) is a common disease that affects 1 to 3 per 1000 individuals per year(1). VTE susceptibility exhibits a complex etiology involving contributions of both genes and environment. Genetic risk factors explain approximately 60% of the overall risk for VTE(2). Recent large-scale genome-wide association studies (GWAS) confirm *ABO*, *F2 F5*, *F11*, *FGG* and *PROCR* as thrombosis susceptibility genes, with only two additional novel loci, TSPAN15, SLC44A2, identified(3-5), leaving the major component of VTE genetic risk still unexplained.

The Factor V Leiden variant $(F5^L)$ is a common inherited risk factor for VTE with an average allele frequency of 3.5% in the European population(6-8). $F5^L$ is estimated to account for up to 25% of the genetically-attributable thrombosis risk in humans(6). However, penetrance is incomplete, with only 10% of $F5^L$ heterozygotes developing thrombosis in their lifetimes. The severity of thrombosis also varies widely among affected individuals(7, 9), limiting the clinical utility of $F5^L$ genotyping in the management of VTE(10).

The incomplete penetrance and variable expressivity of thrombosis among $F5^L$ patients can at least partially be explained by genetic interactions between $F5^L$ and other known thrombotic risk factors such as hemizygosity for antithrombin III or proteins C or S, as well as the common prothrombin 20210 polymorphism(9, 11, 12). However, <2% of $F5^L$ heterozygotes would be expected to co-inherit a mutation at one or more of these loci, suggesting that a large number of additional genetic risk factors for VTE and/or modifiers of $F5^L$ remain to be identified(3, 9).

Mice carrying the orthologous $F5^L$ mutation exhibit a mild to moderate prothrombotic phenotype closely mimicking the human disorder(13). We previously reported a synthetic lethal

interaction between $F5^L$ homozygosity ($F5^{L/L}$) and hemizygosity for tissue factor pathway inhibitor ($Tfpi^{+/-}$)(14). Nearly all mice with this lethal genotype combination ($F5^{L/L}$ $Tfpi^{+/-}$) succumb to widespread, systemic thrombosis in the immediate perinatal period(14).

ENU mutagenesis in mice has been used effectively to identify novel genes involved in a number of biological processes(15, 16). ENU-induced germline mutations transmitted from a mutagenized male mouse (G0) occur at ~1.5 mutations per megabase, at least 50 fold higher than the endogenous background mutation rate(17). Several previous reports have successfully applied an existing phenotype as a sensitizer to identify modifier genes. A dominant suppressor screen in MecP2 deficient mice (Rett syndrome) identified a mutation in squalene epoxidase (Sqle) as a heritable suppressor, resulting in prolonged survival and amelioration of neurologic manifestations(18). Other successful sensitized screens include analysis of mouse mutants predisposed to diabetic nephropathy(19), a screen in Sox10 haploinsufficent mice identifying the Gli3 gene as a modifier of neurochristopathy(20) and identification of a mutation in the c-Myb gene as a dominant modifier for platelet count in Mpl deficient mice (congenital thrombocytopenia)(21). We now report the results of a dominant, sensitized ENU mutagenesis screen for suppressors of $F5^{L/L}$ $Tfpi^{+/-}$ dependent lethal thrombosis.

Results and Discussion

F8 deficiency suppresses F5^{L/L} Tfpi^{+/-} lethality

X-linked hemophilia A results in a moderate to severe bleeding disorder in humans and is caused by mutations in the F8 gene. To test whether the $F5^{L/L}$ $Tfpi^{+/-}$ lethal thrombotic phenotype is suppressible by hemophilia A in mice, triple heterozygous $F5^{L/+}$ $Tfpi^{+/-}$ female mice were generated and crossed to $F5^{L/L}$ male mice (Fig. 1A). One quarter of conceptuses are expected to carry the $F5^{L/L}$ $Tfpi^{+/-}$ genotype, with half of the total expected male conceptuses completely F8

deficient ($F8^{\circ}$). Thus, $1/16^{\text{th}}$ of the overall offspring from this mating are expected to be $F5^{L/L}$ $Tfpi^{+/-}F8^{\circ}$ (males). Similarly, $1/16^{\text{th}}$ of the progeny should be $F5^{L/L}$ $Tfpi^{+/-}F8^{+/-}$ (females). A total of 163 progeny from this cross were genotyped at weaning, resulting in $8 F5^{L/L}$ $Tfpi^{+/-}F8^{\circ}$ male mice observed (and $0 F5^{L/L}$ $Tfpi^{+/-}F8^{+}$, p=0.02) and $2 F5^{L/L}$ $Tfpi^{+/-}F8^{+/-}$ female mice (and $1 F5^{L/L}$ $Tfpi^{+/-}$ $F8^{+/+}$, p=0.9). These results demonstrate that $F5^{L/L}$ $Tfpi^{+/-}$ thrombosis is genetically suppressible by F8 deficiency with nearly complete penetrance in $F8^{\circ}$ male mice and are consistent with human studies demonstrating F8 level as an important VTE risk factor(22).

The F5^{L/L} Tfpi^{+/-} phenotype is suppressed by dominant ENU induced mutations

A sensitized, genome-wide ENU mutagenesis screen for dominant thrombosis suppressor genes was implemented as depicted in Figure 1B. ENU mutagenized G0 $F5^{L/L}$ males were crossed to $F5^{L/L}$ $Tfpi^{+/-}$ females to generate G1 mice, which were screened by genotyping at weaning for $F5^L$ and $Tfpi^{+/-}$. Previously described visible dominant mutants(23), including belly spotting and skeletal abnormalities, were observed in approximately 5.9% of G1 offspring, similar to the ~4.2% rate of observable mutants in previous studies(23). This is consistent with the ~20-30 functionally significant mutations per G1 mouse expected with this ENU mutagenesis protocol(24, 25). One quarter of G1 embryos from this cross are expected to carry the synthetic lethal $F5^{L/L}$ $Tfpi^{+/-}$ genotype. Out of a total of 6,729 G1 mice screened at weaning, the 98 live $F5^{L/L}$ $Tfpi^{+/-}$ mice (45 females, 53 males) represented 4.4% of the 2,210 embryos expected with this genotype. Survival data were collected for 57 of the $F5^{L/L}$ $Tfpi^{+/-}$ G1 mice, with 34 living past 70 days of age (precise dates of death were not available for the remaining 41 mice). No significant sex-specific differences in survival were observed (Fig. 1C).

Heritability for each of the 98 G1 putative suppressor mutants who lived to breeding age (44/98) was evaluated by a progeny test backcross to C57BL/6J (B6) $F5^{L/L}$ mice. The

observation of one or more $F5^{L/L}$ $Tfpi^{+/-}$ offspring surviving to weaning increased the likelihood that a particular Modifier of Factor 5 Leiden (MF5L) line carries a transmissible suppressor mutation. Out of the original 98 surviving F5^{L/L} Tfpi^{+/-} G1 mice, 75 produced no offspring surviving to weaning, either due to infertility or the above mentioned early lethality, with >50% of these mice (37 of 75) exhibiting a grossly runted appearance. Roughly half of the F5^{L/L} Tfpi^{+/-} G1 mice that attained breeding age (23/44) produced 1 or more G2 progeny surviving to weaning, 7 (2 males and 5 females) produced no F5^{L/L} Tfpi^{+/-} G2s, including 4 G1s with 8 or more offspring of other genotypes. Sixteen $F5^{L/L}$ $Tfpi^{+/-}$ G1 mice produced one or more $F5^{L/L}$ Tfpi^{+/-} progeny when bred to B6 F5^{L/L} mice (see Methods). These 16 potential thrombosuppressor mouse lines (designated MF5L1-16) were crossed onto 129S1/SvIMJ (129S1) to generate suppressor lines of genetically informative progeny. The number of total progeny, genotypic distribution, and penetrance of the F5^{L/L} Tfpi^{+/-} mice in each line are listed in Table S1. Within these suppressor lines, mice with the $F5^{L/L}$ Tfpi^{+/-} genotype were ~30% smaller than their $F5^{L/L}$ littermates at the time of weaning (p<2.2x10⁻¹⁶, Fig. 1D), with this difference maintained after outcrossing to the 129S1 strain (Fig. 1E).

Previous reports based on gene function in the specific locus test estimate an ENU-induced mutation rate of 1/700 loss of function mutations per locus for the ENU dosing regimen used here(26). This mutation rate predicts that our screen of 6,729 G1 progeny (2,210 $F5^{L/L}$ $Tfpi^{+/-}$ expected) should have produced ~3 mutations per gene averaged over the entire genome, with 54% of these mutations expected to be null(27), resulting in 1.5X genome coverage for loss of function mutations.

The MF5L6 suppressor mutation maps to a chromosome 3 interval containing F3

In order to map putative ENU-induced suppressor mutations, surviving $F5^{L/L}$ $Tfpi^{+/L}$ mice were intercrossed with $F5^{L/L}$ mice that had been backcrossed onto the 129S1 strain. Crosses between $F5^{L/L}$ and $F5^{L/L}$ $Tfpi^{+/L}$ mice (both $F5^L$ and $Tfpi^-$ backcrossed > 12 generations onto 129S1) confirmed the lethality of the $F5^{L/L}$ $Tfpi^{+/L}$ genotype on the 129S1 background (Table S2).

Each MF5L suppressor line was intercrossed to 129S1 mice. The 4 lines containing the largest number of genetically informative B6-129S1 mixed background $F5^{L/L}$ $Tfpi^{+/-}$ offspring (MF5L1, 6, 9 and 16) were used for gene mapping. Though the MF5L1, MF5L9 and MF5L16 were successfully expanded to pedigrees containing 27, 84, and 14 $F5^{L/L}$ $Tfpi^{+/-}$ informative offspring, respectively, genotyping for a total of ~800 markers in each cross failed to identify any loci with a LOD $\gtrsim 3$ (maximum LODs for MF5L1=1.15, MF5L9=2.5 and MF5L16=1.61). The absence of a clear linkage signal for each of these lines likely reflects complex mouse strain modifier gene interactions, which are known to significantly impact mouse phenotypes(9, 28) and confound linkage analysis(29). Consistent with this hypothesis, we have previously documented poorer survival to weaning in mixed B6-129S1 $F5^{L/L}$ mice compared to littermates(13). We extended these observations by the analysis of additional $F5^{L/L}$ and $F5^{L/L}$ littermates, with mice of the $F5^{L/L}$ genotype demonstrating a 50% reduction in survival in the 129S1 versus B6 strain backgrounds (Table S2).

MF5L6 was maintained for 12 generations on both the mixed and B6 backgrounds and produced a total of 336 $F5^{L/L}$ $Tfpi^{+/-}$ mice (98 on the mixed B6-129S1 background and therefore useful for linkage analysis, See Table S1). Genome-wide SNP genotyping was performed on DNA from these 98 genetically informative $F5^{L/L}$ $Tfpi^{+/-}$ mice, with multipoint linkage analysis shown in Figure 2A. Since the genetic intervals around the F5 and Tfpi loci cannot be accurately assessed for linkage, these regions of chromosomes 1 and 2 were excluded from linkage analysis

(See Fig. 2A Legend and Methods). A single locus with a significant LOD score of 4.49 was identified on chromosome (Chr) 3, with the 1 LOD interval (117.3-124.8Mb) containing 43 refseq annotated genes (Fig. 2B).

The F3 gene located within this interval (Chr3:121.7 Mb) (Fig. 2B) encodes tissue factor (TF), a procoagulant component of the hemostatic pathway that has *Tfpi* as its major regulator. Quantitative or qualitative deficiencies in F3 are thus highly plausible candidates to suppress the $F5^{L/L}$ Tfpi^{+/-} phenotype. To test F3 as a candidate suppressor of the $F5^{L/L}$ Tfpi^{+/-} phenotype, an independent F3 null allele was introduced and triple heterozygous $F5^{L/+}$ $Tfpi^{+/-}$ $F3^{+/-}$ mice were crossed to F5^{L/L} B6 mice (Fig. 2C). Of 273 progeny genotyped at weaning, 13 F5^{L/L} Tfpi^{+/-} F3^{+/-} were observed (and $1 F5^{L/L} Tfpi^{+/-} F3^{+/+}$, p=9.7x10⁻⁵). We also observed significantly fewer male than female $F5^{L/L}$ $Tfpi^{+/-}$ $F3^{+/-}$ mice (2 vs. 11 p=0.03). Thus, haploinsufficiency for $F3^{+/-}$ suppresses the synthetic lethal $F5^{L/L}$ $Tfpi^{+/-}$ phenotype, although with incomplete penetrance (33%) that also differs by sex (10% for males and 67% for females). In contrast, the MF5L6 line exhibited an overall penetrance of 72.4%, with similar male/female penetrance. Gender-specific differences in venous thrombosis rates have previously been reported, including contributions from oral contraceptives and hormone replacement therapy(30-32). This difference in penetrance could be due to 129S1 strain effects in the MF5L6 line or differences between a F3 regulatory mutation in MF5L6 compared to the F3 loss of function allele.

Whole exome sequencing data analysis of a $F5^{L/L}$ $Tfpi^{+/-}$ mouse from MF5L6 failed to identify an ENU variant in F3 or in any other genes in the nonrecombinant interval, or more broadly on the entire Chr3. Although additional ENU variants were identified on other chromosomes, none co-segregated with the survival phenotype in line MF5L6 (Table S3). Sanger sequencing analysis of the full set of F3 exons and introns, as well as 5kb upstream of exon 1,

also failed to identify an ENU-induced mutation. In addition, analysis of *F3* mRNA levels in liver, lung and brain tissues of adult mice failed to identify any differences in the level of expression from the ENU-mutant compared to the wildtype allele (Fig. S1).

Taken together, these data suggest that an ENU-induced F3 regulatory mutation outside of the sequenced segment may be responsible for thrombosuppression in MF5L6, although we cannot exclude a regulatory mutation in another gene. Nonetheless, our findings demonstrate that F3/Tfpi balance plays a key role in thrombosis in the mouse, particularly in the setting of $F5^L$, and suggest that modest variations in either F3 or Tfpi could be important modifiers of VTE susceptibility in humans.

Whole exome sequencing identifies candidate ENU suppressor variants for 8 MF5L lines

Whole exome-next generation sequencing (NGS) was performed on genomic DNA from an index $F5^{L/L}$ $Tfpt^{+/-}$ mouse (from the G2-G5 generation) from each of 8 MF5L lines, including the 4 lines described above, as well as 4 additional lines with large pedigrees (MF5L5, MF5L8, MF5L11, MF5L12). The mean coverage of sequenced exomes was more than 90X, with >97% of the captured region covered with at least 6 independent reads (Table S4). A total of 125 heterozygous variants were identified as candidate suppressor mutations, with 79 variants affecting protein sequence (Table S3). Of the total mutations, 54.4% were nonsynonymous single nucleotide variants (SNVs), followed by UTR (17.6%), synonymous (14.4%) and stopgain SNVs (7.2%), with the remainder being comprised of indels, splicing, and stoploss mutations. The most common mutation events were A/T \rightarrow G/C transitions (35.3%), while C/G \rightarrow G/C transversions were the least represented (2.5%). This spectrum of mutations is consistent with previously published ENU experiments(33). Variants exhibiting no recombination with the Tfpi locus on Chr2 (17 variants) were excluded from further analysis (See Methods). Sanger

sequencing confirmation was performed for 62 variants, including all nonsynonymous and stopgain mutations. These variants were then checked for parent of origin (either the G1 mutagenized progeny or its non-mutagenized mate) as well as the original mutagenized G0 male. Forty-seven of these variants were identified in the G1 mouse but not in the G0 or non-mutagenized parent, consistent with ENU-induced mutations. The remaining 15 mutations were either NGS sequencing errors (11/15), de novo (2/15) or transmitted from the non-mutagenized parent (2/15) (Table S3).

Each SNV was analyzed in additional MF5L mice from the line in which it was identified. None of the thrombosuppressive exonic ENU-induced variants identified in lines MF5L1, 5, 6, 8, 9, 11 and 16 segregated with the lethal phenotype as tested by Kaplan-Meier analysis using a significance threshold of p<0.05(34). Of the 7 candidate ENU-induced SNVs identified from whole exome sequence analysis for the MF5L12 line, 1 was an NGS sequencing error and 6 were validated by Sanger sequencing as consistent with ENU-induced mutations in the G0 mice (Table S3). For each of these 6 SNVs, co-segregation with the survival phenotype was tested by Kaplan-Meier analysis of 31 $F5^{L/L}$ $Tfpi^{+/-}$ mice from the MF5L12 line. Only one variant, a nonsynonymous SNV in the Actr2 gene (c.772C>G, p.R258G, $Actr2^G$), demonstrated a significant survival advantage(23) when co-inherited with the $F5^{L/L}$ $Tfpi^{+/-}$ genotype (p=1.7x10⁻⁶) (Fig. 3A).

Actr2 as a candidate thrombosuppressor gene

The *Actr2* gene encodes the ARP2 protein, which is an essential component of the Arp2/3 complex(35). ARP2, along with ARP3 and five other independent protein subunits (ARPC1-5), forms the evolutionarily conserved seven-subunit Arp2/3 complex(36). Arp2/3 is a major component of the actin cytoskeleton and is found in most eukaryotic cells including platelets(37).

Arp2/3 binds to the sides of actin filaments and initiates the growth of a new filament, leading to the creation of branched actin networks that are important for many cellular processes(38). Loss of Arp2/3 function can have severe consequences as illustrated by the embryonic lethality of mice homozygous for an ARP3 hypomorph(39). In hemostasis, the Arp2/3 complex is necessary for actin-dependent platelet cytoskeletal remodelling events, which are essential for platelet activation and degranulation(40-42). The *Actr2*^{+/G} mutation results in a p.R258G substitution in exon 7 of *Actr2*, at a highly conserved amino acid position, with arginine present at this position for all 60 available vertebrate sequences (https://genome.ucsc.edu), as well as in plants and fungi (Fig. 3B). In addition, no variants at this position have been identified to date in over 120,000 human alleles(43).

Actr2 hemizygosity is incompatible with survival

We attempted to generate an independent *Actr2* knockin (*Actr2*^G) allele by CRISPR/Cas9 genome editing. CRISPR/Cas9-induced homology directed repair (HDR)-mediated DNA integration within the *Actr2* gene was tested in the Neuro-2a (N2a) cell line using two single guide RNAs (sgRNAs) and their respective single-stranded DNA (ssDNA) donor sequences for HDR-mediated knockin (Fig. S2A and B, Table S5). In post-selected heterogenous N2a cells, we observed integration of the variant containing DNA sequence when using sgRNA #1 (Fig. S2C).

We next injected 30 blastocysts and analyzed the 13 surviving to the 60-cell stage. Genotyping of the 13 blastocysts produced from the sgRNA #1 injected oocytes exhibited a high degree of mosaicism, with all 13 positive for DNA cleavage events and 5 (38%) positive for *Actr2*^G substitution (Fig. S3). However, injection of 305 fertilized eggs and subsequent transfer of 275 embryos into 9 foster mothers resulted in the generation of only 18 surviving pups at weaning, all of which were wildtype for the *Actr2* allele. Taken together with the high *Actr2*-

targeting efficiency (see Fig. S3 and Methods) these data suggest that heterozygous loss of function for *Actr2* may be incompatible with survival to term.

Consistent with this hypothesis, human sequencing data from the Exome Aggregation Consortium (ExAC), which includes 60,706 individual exomes, reports a loss of function intolerance for *ACTR2* of 0.997(43). *ACTR2* mutations have not been previously associated with human disease (https://omim.org/entry/604221)(44), again strongly suggestive of early embryonic lethality. In addition, out of 373,692 mouse ENU-induced mutations listed in the Mutagenetix website, only 16 are located in the *Actr2* gene, with no predicted loss of function mutations (https://mutagenetix.utsouthwestern.edu/)(45). Taken together, these data strongly suggest that haploinsufficiency for *Actr2* is not tolerated in humans or mice. The viability of *Actr2*^{+/G} mice suggests that the *Actr2*^G allele is either hypomorphic or a unique gain of function mutation distinct from simple haploinsufficiency. Similarly, analysis of ExAC data suggests that 4 of the 6 other members of the Arp2/3 complex are intolerant to heterozygous loss of function in humans(43). Thus, the high efficiency of CRISPR/Cas9 in generating compound heterozygous loss of function variants together with the less efficient *Actr2*^G substitution likely explains the outcome of our *Actr2* genome editing experiments.

The identification of novel factors involved in the regulation of hemostasis is challenging, as genes leading to marked shifts in hemostatic balance resulting in either severe bleeding or thrombosis are straightforward to identify clinically in humans, whereas subtle shifts are likely to escape detection given the multiple layers of buffering built into the complex hemostatic system(46). The dominant sensitized suppressor screen reported here was undertaken to identify genes for which modest (≤50%) reduction in function would significantly shift overall hemostatic balance. Such loci represent likely candidates for common human variation

contributing to thrombosis and bleeding disorders. Gene variants with subtle yet significant antithrombotic effects represent attractive therapeutic targets because of a potentially wide therapeutic window with few unintended side effects. In contrast, a recessive screen would have been expected to identify known Mendelian bleeding disorders such as hemophilia A (F8 deficiency), as confirmed by the data in Figure 1A. The finding of $98 F5^{L/L} Tfpi^{+/-}$ mice carrying putative thrombosis suppressor mutations (at an estimated 1.5X genome coverage) suggests that subtle alterations at a number of loci are capable of suppressing the $F5^{L/L} Tfpi^{+/-}$ lethal thrombotic phenotype. The complex strain-specific genetic modifiers that confounded the genetic linkage analysis are consistent with this model. Nonetheless, our findings illustrate the particular importance of the F3/Tfpi axis in thrombosis regulation (especially in the setting of $F5^L$), as well as the identification of Actr2 and the Arp2/3 complex as another potentially sensitive regulatory pathway for maintaining hemostatic balance.

Materials and methods

Mice

C57BL/6J (B6, stock number 000664), DBA2/J (DBA, stock number 000671), 129S1/SvImJ mice (129S1, stock number 002448), and B6D2F1 mice (stock number 100006) were purchased from the Jackson Laboratory. $F5^{L/L}$ ($F5^{tm2Dgi}$ /J stock number 004080) mice were previously generated(22). F3 and Tfpi deficient mice were a generous gift of Dr. George Broze(47, 48). F8 deficient mice were a generous gift of Dr. Haig Kazazian(49). All mice designated to be on the B6 background were backcrossed greater than 8 generations to B6. $F5^{L/L}$ breeding stock for the 129S1 modifier gene crosses and genetic mapping experiments were generated from $F5^L$ mice serially backcrossed greater than 12 generations to the 129S1 strain to create B6-129S1 $F5^L$ congenic mice. $Tfp\bar{i}$ breeding stock for the 129S1 modifier gene crosses

were generated by serially backcrossed greater than 12 generations to the 129S1 strain to create B6-129S1 $Tfpi^{+/-}$ congenic mice. G1 suppressor mutant heritability was evaluated by a progeny test backcross to B6 $F5^{L/L}$ mice. Among the 16 mice able to produce surviving $F5^{L/L}$ $Tfpi^{+/-}$ offspring, males were overrepresented (4 female, 12 male), likely because of larger numbers of offspring resulting from breeding to multiple female partners. These 16 potential thrombosuppressor mouse lines were crossed onto 129S1 to generate suppressor lines of genetically informative progeny for genetic mapping. The University of Michigan Institutional Committee on the Use and Care of Animals (IACUC) approved all experiments using mice (protocol numbers PRO00007371, PRO00005191 and PRO00005913).

Genotyping

DNA was isolated from tail biopsies and mice were genotyped for $Tfpi^{+/-}$ and $F5^L$ as previously described(14). Mice were genotyped for F3 deficiency using custom primers listed in Table S6. All primers were purchased from Integrated DNA Technologies (IDT), Coralville, IA.

ENU mutagenesis and breeding

ENU was purchased (Sigma Aldrich, St. Louis MO) in ISOPAC vials, and prepared according to the following protocol: http://pga.jax.org/enu_protocol.html. A single ENU dose of 150 mg/kg was administered intraperitoneally into an initial cohort of 159 $F5^{L/L}$ B6 male mice (referred to as generation 0 or G0 mice). For a second cohort of 900 male $F5^{L/L}$ G0 mice, the protocol was changed to three weekly intraperitoneal injections of ENU (90 mg/kg). After a 10-week recovery period, each G0 mouse was bred to $F5^{L/+}$ $Tfpi^{+/-}$ mice (Fig. 1B) on the B6 genetic background to produce G1 generation offspring, which were genotyped at two weeks of age. G1 mice of the $F5^{L/L}$ $Tfpi^{+/-}$ genotype surviving to weaning age (three weeks of age) were considered to carry a suppressor mutation.

Modifier gene transmission

 $F5^{L/L}$ $Tfpi^{+/-}$ G1 founders were crossed to $F5^{L/L}$ mice on the B6 genetic background to produce G2 generation offspring. G2 mice were outcrossed to $F5^{L/L}$ mice on the 129S1 genetic background for 2 or more generations.

Genetic mapping

Genetic markers distinguishing the B6 and 129S1 strains distributed across the genome were genotyped using the Illumina GoldenGate Genotyping Universal-32 platform (Illumina, San Diego CA) at the University of Michigan DNA Sequencing Core. Linkage Analysis was performed on the Mendel platform version 14.0(50) using 806 informative markers from the total of 1449 genotyped markers. LOD scores >=3.3 were considered significant(51). Chrs 1 and 2 contained the *F5* and *Tfpi* genes, respectively, and therefore these chromosomes were excluded from further analysis. The number of mice and the LOD scores for each of the mapped pedigrees are listed in Table S1.

Sanger sequencing of the F3 gene and analysis of candidate mutations

Genomic DNA was extracted from mouse tail biopsies using the Gentra Puregene Tissue Kit (Qiagen, Germantown, MD). A total of 48 overlapping pairs of amplicons (primers: F3gene_1-F3gene_35; upstreamF3_1-upstreamF3_13, Table S6) were used to Sanger sequence the entire F3 gene (~11kb) and an additional ~5kb of upstream sequences on both strands. Sanger sequencing was performed at the University of Michigan Sequencing Core. For the analysis of candidate mutations, amplicons were generated harboring the nucleotide of interest using a custom outer primer pair. Inner forward and reverse primers were used to bidirectionally sequence these amplicons. Sequencing chromatograms were visualized and manually scored using FinchTV (PerkinElmer, Waltham, MA).

Estimation of F3 allelic expression

The F3 exonic region harbors 3 known B6-129S1/DBA SNPs (rs30268372, rs30269285, rs30269288, http://www.ncbi.nlm.nih.gov/SNP/) that were used for relative expression analysis. F5^{L/L} Tfpi^{+/-} mice with one B6 allele (in cis with ENU induced variants) and one 129S1 allele at the Chr3 candidate region were outcrossed to DBA wildtype females introducing exonic B6-129S1/DBA SNPs. Five progeny from this cross (2 B6/DBA and 3 129S1/DBA allele carriers, identified by DNA genotyping) were tested for differential allelic expression. Three tissue samples (lung, liver, whole brain) were obtained from each mouse as previously described(22). RNA was extracted from the tissue samples using RNeasy Plus Mini Kit (Qiagen) according to manufacturer's recommendations and reverse transcribed using SuperScript II (Invitrogen, Carlsbad, CA). cDNA corresponding to exon 3-exon 5 was amplified with primers F3-exon-F and F3-exon-R using GoTaq Green Master Mix (Promega, Madison, WI). Primers F3-exon-F and F3-exon-R were also used to Sanger sequence the F3 exonic region.

Relative expression was estimated at SNP sites by dividing the area under the Sanger sequencing peak of one allele to another (52, 53). Next, the relative expression of each SNP was compared between the B6 and 129S1 allele carrying progeny.

Mouse whole-exome sequencing

Libraries were prepared using Agilent (Agilent Technologies, Santa Clara, CA) or NimbleGen (Roche NimbleGen, Madison, WI) mouse whole exome capture kits. 100 base pair (bp) paired-end sequencing was performed on the Illumina Hiseq 2000 platform at the University of Michigan DNA Sequencing Core. A detailed overview of the whole exome sequencing pipeline is available at GitHub (https://github.com/tombergk/FVL_SUP). Briefly, sequence reads were aligned using Burrows-Wheeler Alignment software(54) to the mouse reference genome

(genome assembly GRCm38, Ensembl release 73). Reads were sorted and duplications removed using Picard tools (http://picard.sourceforge.net). Coverage statistics were estimated using QualiMap software(55). Variants were called across 8 samples using GATK HaplotypeCaller software(56). Standard hard filters recommended by the Broad Institute were applied using GATK VariantFiltration(56) followed by an in-house developed pipeline to remove variants between the B6 and 129S1 strains, shared variants within our mouse cohort and variants in closer proximity than 200 bp from each other. Variants were annotated using Annovar software(57) with refseq annotation (release 61). Heterozygous variants within exonic regions with >6X coverage unique for only one mouse in the cohort were regarded as potential ENU induced variants. A total of 125 heterozygous variants were identified as candidate suppressor mutations, using an in-house filtering pipeline(52), with 79 variants occurring within coding sequence. The number of ENU variants identified in each exome sequenced mouse varied by genealogical distance from the G1 *MF5L* founder. The candidate ENU induced variants were confirmed by Sanger sequencing.

Generation of Actr2 CRISPR/Cas9 targeted mice and cells

Actr2 Targeting Sequence Design and Cloning: The pX459 (pSpCas9(BB)-2A-Puro; plasmid ID:48139)(58) biscistronic expression vector for human-codon optimized *S. pyogenes* Cas9, chimeric sgRNA and puromycin resistance gene was obtained from Addgene. The vector was digested with *BbsI* and a pair of annealed oligos (sgRNA) was independently cloned into the backbone vector as described(58) and depicted in Figure S2A. Colonies were screened for successful guide insertion by *BbsI/AgeI* double digestion to discriminate between positive and negative clones. Clones that displayed only a ~8.5 kb linearized *AgeI* digested fragment were used in downstream applications.

The *Actr2*-specific sgRNA sequences used in these experiments are listed in Table S5. sgRNAs were selected based on: 1) their proximity to the mutation site (< 100 bp away; 10 bp optimal); 2) the presence of a protospacer adjacent motif (PAM) "NGG" sequence adjacent to the sgRNA; 3) the ability to incorporate a synonymous variant within the PAM to protect the HDR donor template from Cas9-targeted degradation; and 4) the sequence must have an inverse likelihood of off-target binding score of >70 and no less than 3 mismatches within exonic regions of the genome as determined by the CRISPR design tool (crispr.mit.edu).

Single-Stranded DNA Donor for Homology Directed Repair: The ssDNA oligonucleotides that served as HDR donor templates were ordered as Ultramer DNA oligos from IDT and are listed in Table S5. The HDR donor template consists of a 161 bp genomic sequence homologous to a region spanning -44 to +117 bp from the splice junction of intron 6 and exon 7 of the mouse Actr2 gene (Fig. S2B). The HDR donor encodes an arginine (R) to glycine (G) mutation at the 258th amino acid position (c.772C>G) and a synonymous mutation within the PAM to prevent donor DNA cleavage by Cas9. In this design, 80 bp homology arms flank the C to G transversion mutation with the position of the double-strand break (DSB) occurring 11 – 12 bp downstream or 17 – 18 bp upstream of the homology arm junction for sgRNA #1 and #2 respectively.

Transfection of Neuro-2a cells for Validation of sgRNA and HDR Efficiency: Mouse N2a cells (ATCC® CCL-131TM) were routinely cultured in EMEM (Hyclone, Logan, UT), supplemented with 10% (v/v) fetal bovine serum (FBS) (Hyclone) and incubated at 37°C in the presence of 5% CO2. N2a cells (passage 4) were seeded in triplicate into 24-well plates at a density of 1 x 10⁵ cells per well in 0.5 ml EMEM containing 10% FBS 24 h prior to transfection. Cells in each well were co-transfected for 24 h with 0.5 μg pX459 plasmid (expressing either sgRNA #1 or #2) and 0.5 μg HDR donor template using 0.75 μl lipofectamine 3000 (Invitrogen/Thermo Fisher

Scientific, Waltham, MA) diluted in Opti-Mem I media (Gibco/Thermo Fisher Scientific) following manufacturer's instruction. Transfected cells were isolated by antibiotic selection using 2 µg/ml Puromycin Dihydrochloride (Gibco/Thermo Fisher Scientific) for 72 h. Transfected cells were passaged once prior to harvesting for genomic DNA extraction using the Purelink Genomic DNA mini kit (Invitrogen/Thermo Fisher Scientific) according to manufacturer's protocol.

Mouse Pronuclear Injection: Pronuclear microinjection was carried out essentially as described in Brinster et al. (59). Following pronuclear microinjection mouse zygotes were cultured *in vitro* to the blastocyst stage prior to DNA extraction and analysis for the presence of indels and mutations. Establishing stable CRISPR mouse lines by pronuclear co-injection of Cas9 mRNA, sgRNA and oligo donors is highly efficient, as it was reported that for every 100 embryos that underwent this process, ~13 genetically modified embryos were produced(60). CRISPR/Cas9 Actr2-edited embryos and mice were generated in collaboration with the University of Michigan Transgenic Animal Model Core (TAMC). A premixed solution of 5 ng/μl of pX459 plasmid containing sgRNA #1 targeting Actr2 exon 7 and 10 ng/μl HDR donor template was prepared in RNAse-free microinjection buffer (10 mM Tris-HCl [pH 7.4], 0.25 mM EDTA) and microinjected into the male pronucleus of fertilized mouse eggs obtained from the mating of B6 male mice with superovulated B6 female mice. Microinjected eggs were transferred to pseudopregnant B6DF1 female mice.

Genome Extraction from Blastocyst Embryos: Mouse blastocyst DNA extraction was performed based on the method described by Sakurai et al.(61) Briefly, 60-cell expanded blastocysts cultured for 3 days in vitro at the TAMC were individually collected into 0.2 ml tubes in 10 μl of ultrapure water. Ten microliters of 2X blastocyst extraction buffer (100 mM)

Tris-HCl [pH 8.3], 100mM KCl, 0.02% gelatin, 0.45% Tween-20 supplemented with 60 μg/ml yeast tRNA and 125 μg/ml Proteinase K) was added and the samples were incubated at 56°C for 10 min followed by 95°C for 10 min and immediately placed on ice to prevent heteroduplex formation. Crude lysates were stored at -20°C.

Multiplex PCR Genotyping: We designed a two-reaction multiplex PCR strategy sensitive enough to detect wildtype (WT) and R258G alleles that differ by a single nucleotide. Common ACTR2_OF forward and ACTR2_OR reverse primers were used in separate reactions with the WT-specific reverse primer (ACTR2_WT-R) or the R258G-specific reverse primer (ACTR2_MUT-R) that only differ in the -1 position on the 3'-end (Table S6). Inclusion of the common primers (659 bp product) provided amplification competition and acted as a positive PCR reaction control. Amplification of the 318 bp product using ACTR2_WT-R or ACTR2_MUT-R allele-specific primers indicates the presence of the wildtype or mutant allele respectively (Figs. S2D and S3A).

SURVEYOR Nuclease Assay: The genomic region flanking the CRISPR target site for *Actr2* was PCR amplified using genomic primers ACTR2_OF and ACTR2_OR (Table S6). Unpurified PCR products (30 μl) were subjected to a denaturing and reannealing process to enable heteroduplex formation: 95°C for 10 min; 95°C to 85°C ramping at -2°C/s; 85°C to 25°C at -0.3°C/s; 25°C for 1 min and 4°C hold. After reannealing, ~250 ng DNA products were treated with SURVEYOR nuclease and SURVEYOR enhancer S (IDT) following the manufacturer's recommended protocol for GoTaq DNA polymerase amplified products. Digested and undigested (Cut/Uncut) products were analyzed by standard gel electrophoresis using 2.0% TAE agarose gels containing ethidium bromide and were imaged with a Chemi-Doc Touch gel imaging system (Bio-Rad).

Statistical Data Analysis

Statistical differences among the potential progeny of mouse crosses were determined using the Fisher's Exact test. The student's t-test was used for estimating statistical differences between the weights of $F5^{L/L}$ $Tfpi^{+/-}$ mice and their littermates. Relative expression differences for F3 alleles were estimated using the Wilcoxon rank-sum test. All the above statistical analyses were performed using the 'stats' package in R software. Kaplan-Meier survival curves with log-rank test to estimate significant differences in mouse survival as well as significance for putative suppressors identified by exome sequencing were performed using the 'survival' package in R(62).

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Figure legends

Figure 1: F8 deficient thrombosuppression and design of the Leiden ENU mutagenesis screen. A. The mating scheme and observed distributions of the $F5^{L/+}$ $Tfpi^{+/-}$ F8 deficiency rescue experiments. F8 X results in incompletely penetrant suppression of the $F5^{L/L}$ $Tfpi^{+/-}$ phenotype. B. The mating scheme and observed distribution of the Leiden screen. $F5^{L/L}$ $Tfpi^{+/-}$ male mice were mutagenized with either 1 x 150mg/kg or 3 x 90 mg/kg ENU and bred with non-mutagenized $F5^{L/+}$ $Tfpi^{+/-}$ females. Sixteen and 83 $F5^{L/L}$ $Tfpi^{+/-}$ progeny, respectively were observed in each of the dosing regimens, with over twice the rate of $F5^{L/L}$ $Tfpi^{+/-}$ survivors in the progeny of the 3 x 90 mg/kg treated mice. C. There was no significant difference in survival between male and female $F5^{L/L}$ $Tfpi^{+/-}$ putative suppressor mice, (p=0.384). D. $F5^{L/L}$ $Tfpi^{+/-}$ putative suppressor mice were significantly smaller than their non- $F5^{L/L}$ $Tfpi^{+/-}$ littermates. E. $F5^{L/L}$ $Tfpi^{+/-}$ putative suppressors were smaller regardless of whether they were on the pure B6 or mixed B6-129S1 genetic backgrounds (p=0.327).

Figure 2: **The** *MF5L6* **suppressor locus maps to Chr3. A.** Linkage analysis for the *MF5L6* line. Alternating red and black is used to highlight the chromosomes. Chrs 1 and 2 contained the *F5* and *Tfpi* genes, respectively, and therefore these chromosomes were excluded from further analysis. The Chr3 peak had the highest LOD score in the Chr3 subregion:117.3-124.8Mb (maximum LOD=4.49, 1 LOD interval, significance threshold of LOD >3.3(63)). **B.** The Chr3 candidate interval (Chr3:117.3-124.8 Mb) contains 43 refseq annotated genes, including *F3*. **C.** The mating scheme and observed distribution of offspring to test *F3* deficiency as a suppressor of $F5^{L/L}$ $Tfpi^{+/-}$. $F3^{+/-}$ results in incompletely penetrant suppression of the $F5^{L/L}$ $Tfpi^{+/-}$ phenotype.

Figure 3: **Discovery and validation of** *Actr2* **R258G** as a thrombosis suppressor gene by **NGS. A.** Kaplan-Meier survival plot for $F5^{L/L}$ $Tfpi^{+/-}$ mice with and without the $Actr2^G$ mutation. $F5^{L/L}$ $Tfpi^{+/-}$ $Actr2^{+/G}$ mice exhibit significantly better survival than $F5^{L/L}$ $Tfpi^{+/-}$ $Actr2^{+/+}$ littermates (n = 35). **B.** ARP2 amino acid R258 is highly conserved in animals, plants and fungi.

Figure 1

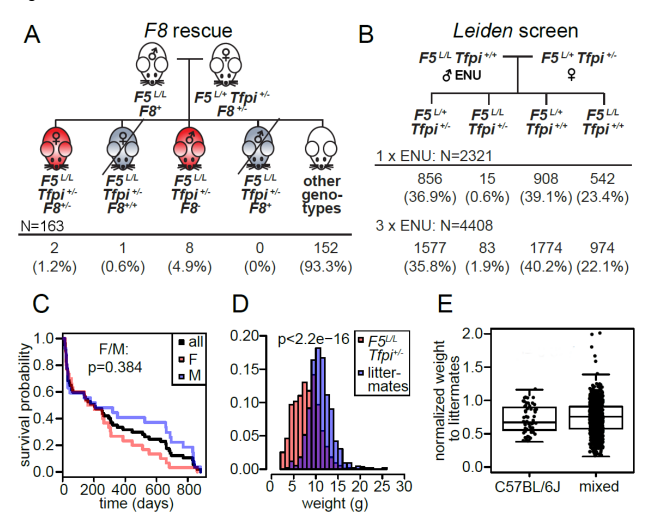


Figure 2

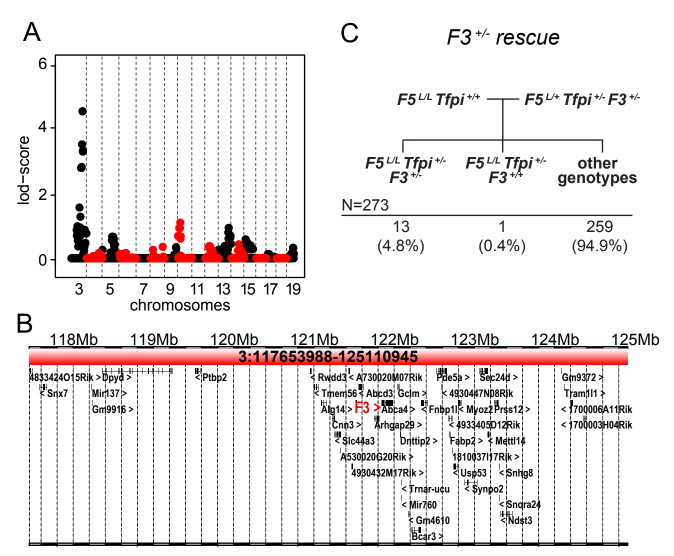


Figure 3

