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Modeling human RNA spliceosome mutations in the mouse: not all mice were created equal

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Myelodysplastic syndromes (MDS) and related myelodysplastic/myeloproliferative neoplasms (MDS/MPNs) are clonal stem cell disorders, primarily affecting patients over 65 years of age. Mapping of the MDS and MDS/MPN genome identified recurrent heterozygous mutations in the RNA splicing machinery, with the *SF3B1*, *SRSF2*, and *U2AF1* genes being frequently mutated. To better understand how spliceosomal mutations contribute to MDS pathogenesis in vivo, numerous groups have sought to establish conditional murine models of *SF3B1*, *SRSF2*, and *U2AF1* mutations. The high degree of conservation of hematopoiesis between mice and human and the well-established phenotyping and genetic modification approaches make murine models an effective tool with which to study how a gene mutation contributes to disease pathogenesis. The murine models of spliceosomal mutations described to date recapitulate human MDS or MDS/MPN to varying extents. Reasons for the differences in phenotypes reported between alleles of the same mutation are varied, but the nature of the genetic modification itself and subsequent analysis methods are important to consider. In this review, we summarize recently reported murine models of *SF3B1*, *SRSF2*, and *U2AF1* mutations, with a particular focus on the genetically engineered modifications underlying the models and the experimental approaches applied. © 2018 ISEH – Society for Hematology and Stem Cells. Published by Elsevier Inc. All rights reserved.

Myelodysplastic syndromes (MDS), myelodysplastic/myeloproliferative neoplasms (MDS/MPN), and related disorders are a heterogeneous class of blood cancers leading to ineffective hematopoiesis in the bone marrow (BM) [1,2]. Approximately 30% of MDS patients progress to acute leukemia. Median survival ranges from 97 months for low-risk categories down to 11 months for high-risk MDS [2]. The incidence of MDS in the general population is approximately four to five per 100,000 people, but this increases with age [1]. Population-based studies in both Australia and the United States indicate a significant underestimation of the true burden of MDS, with frequencies estimated at 103 per 100,000 and between 75 and 162 per 100,000, respectively, over age 65 [3–5]. A feature of MDS and

MDS/MPN is the progressive establishment of clonal hematopoiesis, in which mutant cells dominate the BM at the expense of normal hematopoiesis [6]. The mechanisms behind this clonal advantage are unknown, but the identification of clonal hematopoiesis of indeterminate potential (CHIP) suggests that the clonal advantage is acquired progressively over many years following the establishment of an initiating mutation [7,8]. In general, MDS patients have limited treatment options. Current treatment strategies for MDS and MDS/MPN are mostly heterogeneous and modestly efficacious and are not associated with durable responses [9]. The dismal prognosis following hypomethylating agent failure highlights the urgent need for new treatment approaches.

Until recently, there were few described recurrent mutations or familial syndromes that could provide insight into the genetics of these cancers that could be used to develop preclinical models. Now, however, the

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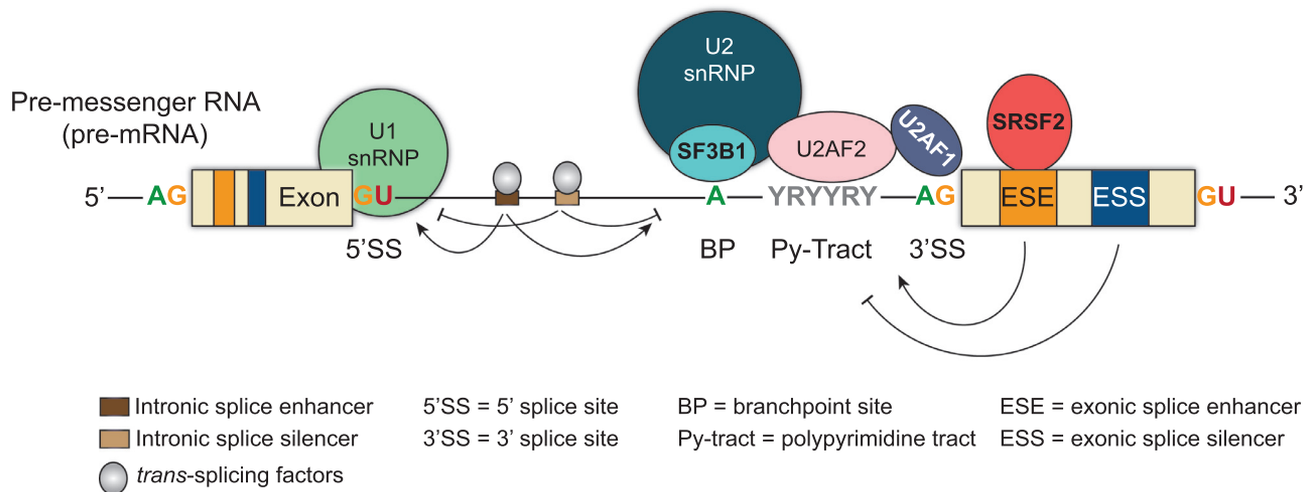


Figure 1. Schematic outline of RNA splicing and the role of those proteins mutated in MDS. Within the intron of the pre-mRNA the U2 snRNP complex containing SF3B1 binds the branch-point site adenosine, U2AF2 binds to the polypyrimidine tract sequence, and U2AF1 binds the 3' splice site; SRSF2 binds to the exonic splice enhancer sequence within the exonic sequence. *Cis*- and *trans*-acting factors can both positively or negative regulate splicing. Proteins in which murine models have been generated are shown in bold font.

detailed mutational architecture of human MDS, chronic myelomonocytic leukemia (CMML), and other related forms of MDS/MPN has been defined. There are recurrent mutations in key pathways, including the RNA-splicing machinery (e.g., *SF3B1*, *SRSF2*, and *U2AF1*) [10–12], transcription factors (e.g., *RUNX1*, *BCOR*, and *ETV6*), and epigenetic enzymes (e.g., *DNMT3A*, *TET2*, *EZH2*, and *ASXL1*) [6]. Additional studies have deduced clonal evolution and mutational order, highlighting distinct patterns [13]. Paralleling studies of diseased samples, analysis of healthy elderly populations has demonstrated that mutations associated with MDS and leukemia accumulate and are positively selected for in the hematopoietic cells from otherwise healthy people ≥ 70 years of age [7,8,14].

The identification of recurrent mutations in the RNA-splicing machinery was a breakthrough in understanding the genetics of MDS and related disorders [10–12]. RNA splicing is a highly coordinated and essential process carried out by major and minor spliceosomes to remove noncoding regions (introns) of the pre-mRNA before protein translation [15,16]. The major spliceosome consists of five small nuclear ribonucleoprotein complexes (snRNPs), U1, U2, U4, U5, and U6. The minor spliceosome includes the U5 snRNPs and other functional snRNPs mirroring the major spliceosome. Mechanistically, RNA splicing occurs with the initial recognition of the 5' and 3' splice sites by the U1 and U2 complexes, respectively, followed by the excision of the intron and exon ligation by U4, U5, and U6 complexes. *Trans*-acting splicing factors, such as serine-arginine-rich proteins and heterogeneous nuclear RNPs, bind to the regulatory elements located in the exons and introns to enhance

or repress the splicing activity and contribute to alternative splicing (Figure 1). The regulatory elements include exonic splicing enhancers and silencers and intronic splicing enhancers and silencers [16].

In MDS, >80% of patients have a mutation in a spliceosome gene [10–12]. In phenotypically overlapping syndromes of MDS/MPN such as CMML, spliceosome mutations are also common [10–12,17]. Mutations occur most frequently in *SF3B1* and *SRSF2*, with a lower prevalence in other spliceosomal genes, including *U2AF1*, *ZRSR2*, and *U2AF2* [17,18]. Spliceosomal mutations are thought to arise early in the course of disease, including as founder/initiating events [6]. However, how mutations in the spliceosome alter normal hematopoiesis and contribute to disease pathogenesis remains unclear. A key experimental system to understand the impact of the spliceosome mutations in regulating both normal and malignant hematopoiesis is the generation of high-fidelity murine models. There is a very high conservation of the human and mouse core spliceosome proteins, including those mutated in MDS and CMML. For example, human *SRSF2* and murine *Srsf2* proteins are 100% sequence conserved.

The generation of conventional null alleles of RNA-splicing genes in the mouse has been useful in identifying key RNA substrates with splicing that is perturbed in their absence and in beginning to understand their normal physiological function [19–22]. Retroviral overexpression has been used to assess how the mutations present in patients may contribute to MDS development [10,22]. A caveat with both the loss-of-function alleles and retroviral overexpression of mutant cDNAs is that they fail to recapitulate the mutations as they occur in humans: heterozygous point mutations

expressed from their endogenous locus, not alleles resulting in protein deficiency. The recurrent finding that the spliceosome mutations are heterozygous indicates that retention of a wild-type (WT) copy is necessary for cells to survive, an interpretation supported by both genetic and pharmacologic evidence [23–26], and additionally that gene dosage and protein complex stoichiometry may be important in the pathogenicity of the mutations [27].

Over the last 3 years, murine models recapitulating point mutations identified in patients with MDS and CMML have been described for *Sf3b1* [28,29], *Srsf2* [30–32], and *U2af1* [33,34]. The initial *U2af1* mutation model reported was an inducible cDNA expressed from a heterologous locus, but will be discussed alongside the true knock-in alleles [33]. The phenotypes observed in these “humanized” models have ranged from very mild through to the development of MDS/MPN. The reasons for these variable phenotypes have been not been systematically outlined, but the targeting strategy itself and the analysis methods may contribute significantly to the understanding of these models. Here, we discuss in detail the murine models of these mutations developed to date, their phenotypes and how they compare to the phenotypes of humans with these mutations.

Murine models of RNA spliceosome mutations

U2AF1 (previously known as *U2AF35*)

U2 small nuclear RNA auxiliary factor 1 (*U2AF1*) binds to the AG nucleotides and polypyrimidine tract located near the 3′ splice site in the intronic sequence (Figure 1). It then recruits *U2AF2* and the *U2* major spliceosome complex to facilitate 3′ splicing site recognition [15,35]. *U2AF1* mutations have been identified in lung, breast, colon, and various epithelial carcinomas [36]. In MDS, approximately 11% of patients have a *U2AF1* mutation, predominantly at serine 34 (S34F/S34Y) and glutamine 157 (Q157R/Q157R) [10,37]. One transgenic and two knock-in models have been developed to date.

U2AF1^{S34F} transgenic model (Shirai et al. [33]).

Shirai et al. [33] reported the generation of a transgenic model of *U2AF1* mutation. In this case, unlike the others that will be discussed, an inducible (tetO) human *U2AF1*^{S34F} or control *U2AF1*^{WT} cDNA were targeted to the *Collagen1a1* (*Coll1a1*) locus of *Rosa26-M2rtTA* ES cells [38]. Expression of the WT or S34F cDNAs was not regulated in the manner of the endogenous *U2af1* locus. To activate the expression of the *U2AF1*^{S34F} or *U2AF1*^{WT} cDNAs, animals were administered doxycycline, enabling dose-dependent induction of expression of the human *U2AF1* cDNAs in the

presence of two endogenous wild-type gene copies derived from the mouse (Figure 2).

No analysis was reported of the effects of *U2AF1*^{S34F} on native hematopoiesis (i.e., in animals treated with doxycycline in the absence of prior bone marrow transplantation). The only data presented were from irradiated transplantation recipients reconstituted with whole BM of *U2AF1*^{S34F} M2rtTA or *U2AF1*^{WT} M2rtTA donors. Once reconstitution was established, the recipient animals were administered doxycycline [33]. A similar approach was chosen in a second report of this model [26]. The use of BM transplantation prior to gene induction restricts expression of the mutation to the reconstituted hematopoietic cells, of particular importance when using an inducer expressed from the widely expressed *Rosa26* locus or others such as *Mx1* that are active in non-hematopoietic cell populations [39–41]. A caveat of analysis in the post-transplantation setting is that BM transplantation is known to induce hematopoietic stress, permanently modifying the BM microenvironment [42], and to alter the clonal dynamics of hematopoietic stem cells (HSCs) [43].

Using this experimental approach, the authors showed that the doxycycline-treated *U2AF1*^{S34F} M2rtTA recipients developed a stable peripheral blood leukopenia without changes in red blood cell or platelet numbers for up to 12 months. The leukopenia was the result of reduced B cells and monocytes and was dependent on the continued expression of the mutant *U2AF1*. Within the BM, there was evidence of increased cell death, reduced B cells and monocytes, and increased neutrophils. There was an increase in the percentages of common myeloid progenitors and a subtle increase in the percentage of lineage⁻/c-Kit⁺/Sca-1⁺ (LKS) cells. In competitive transplantations, *U2AF1*^{S34F} cells were serially transplantable through to tertiary recipients, albeit with a reduced chimerism. With extended aging (up to 500 days after BM transplantation), the *U2AF1*^{S34F}-expressing mice were not reported to develop dysplasia, MDS, or acute myeloid leukemia (AML).

***U2af1*^{S34F} knock-in models.** Fei et al. [34] reported the generation of two conditional knock-in models of *U2af1*^{S34F} mutation targeted to the endogenous locus. The S34F mutation is encoded by exon 2 of the murine *U2af1* allele. Both models required Cre protein to induce expression of the mutant allele. Two different knock-in approaches were described, one using an inverted exon 2 (referred to in that study as *IES34F*) and the second using a minigene (referred to as *MGS34F*), which resulted in significant differences in the expression of the mutant allele (Figure 2). Both were tested as heterozygous alleles, one of which encoded endogenous wild-type *U2af1* and the second the mutant allele.

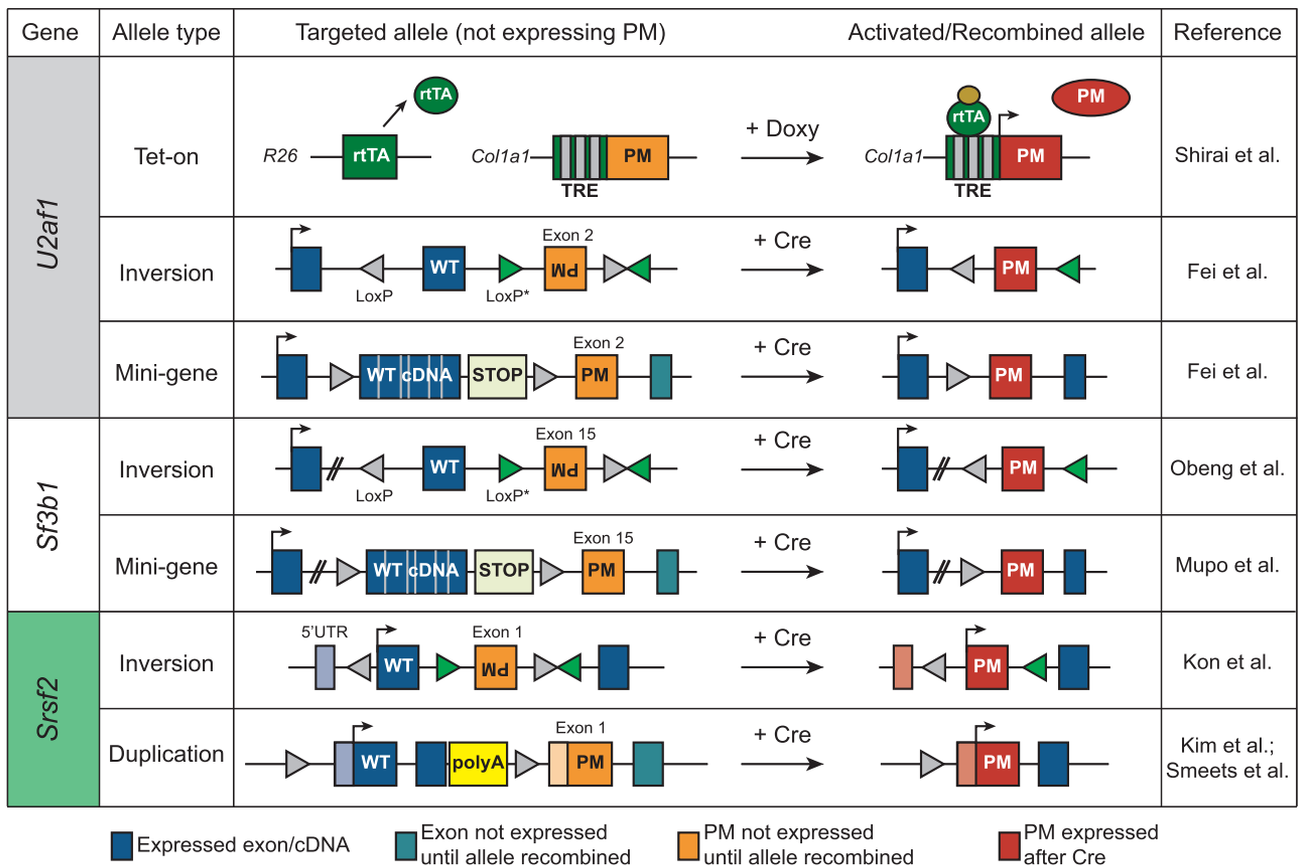


Figure 2. Simplified schematic outline of the currently described spliceosome mutant mouse models. LoxP elements are indicated as grey or green triangles indicating different types of loxP elements. Expressed exons are shown in blue, exons not expressed until after locus recombination are shown in teal, and mutation-bearing exons are shown in orange before recombination (silent) and in red when recombined and expressed.

U2af1^{S34F} “inverted exon” model (Fei et al. [34]).

For the inverted exon 2 model, a lox2272-flanked inverted S34F encoding exon 2 was placed in intron 2–3 and a conventional LoxP element was placed 5' of the endogenous exon 2. Cre-induced recombination of the different LoxP sequences enabled removal of the endogenous exon 2 and replacement by an in-frame exon 2 encoding the S34F mutation. Analysis of mutant allele expression after Cre recombination in mouse embryonic fibroblasts demonstrated that the mutant mRNA was expressed at ~30% despite being genomically heterozygous. It is not clear why the *IES34F* allele failed to express the transcript heterozygously, although the authors suggested that it related to the efficiency of recombination of the locus. This allele was not further characterized [34].

U2af1^{S34F} “minigene” model (Fei et al. [34]). The minigene model was examined in more detail [34]. In this model, a LoxP-flanked cDNA sequence encoding exons 2–8 followed by a transcriptional stop sequence was placed in the first intron. The S34F mutation was inserted in the endogenous exon 2. In the absence of

Cre, the *U2af1* transcript from the targeted allele should be encoded by the endogenous exon 1 together with the cDNA encoding exons 2–8, with the stop cassette preventing read-through and expression from the S34F-encoding allele. This model was tested in the setting of native hematopoiesis using the broadly expressed *Mx1-Cre* and polyI:polyC (pI:pC) induction. The analysis of recombination following pI:pC administration indicated that the minigene, unlike the inverted exon model, was efficiently recombined and truly heterozygous both genomically and transcriptionally. Analysis of the *U2af1*^{S34F/+} animals after pI:pC demonstrated a mild persistent macrocytic anemia and a twofold reduction in leukocytes, particularly B cells, in the peripheral blood. Within the BM, total cellularity was preserved, but there were reductions in the percentages of LKS⁺ and LKS⁺CD48⁻CD150⁺ phenotypic HSCs. Lineage distribution of more committed populations within the BM was not described. Morphological analysis of the BM revealed that less than 1% of cells displayed dysplastic features and no dysplastic cells were reported in the peripheral blood. No MDS or

AML was observed in aged *U2af1*^{S34F/+} mice. The phenotypes described for native hematopoiesis were preserved, albeit exacerbated, by BM transplantation where expression of *U2af1*^{S34F/+} was activated after transplantation, demonstrating a cell-autonomous effect of the mutant allele on hematopoiesis. Interestingly, and consistent across the splicing mutation models reported to date, there was a significantly reduced repopulating potential of *U2af1*(S34F)-expressing cells when BM cells from donors previously treated with pI:pC competed against WT BM cells.

SF3B1. Splicing factor 3b subunit 1 (SF3B1) is part of the SF3b protein complex that recognizes the branch point adenosine (A) base within the intron and facilitates the binding of U2 snRNP on pre-mRNA (Figure 1). *SF3B1* mutations have been identified in MDS, chronic lymphocytic leukemia and uveal melanoma. Mutations cluster between exons 14 and 16, with K700E being the most frequent mutation [10,11]. In MDS, *SF3B1* mutations occur in >80% of refractory anemia with ring sideroblasts, an MDS subtype with a favorable outcome [11,44]. Two independently generated *Sf3b1* conditional knock-in models have been reported and both have utilized the *Mx1*-Cre system to induce the expression of the mutant allele.

Sf3b1^{K700E} knock-in models

Sf3b1^{K700E} “inverted exon” model (Obeng et al. [28]).

A conditional *Sf3b1*^{K700E} model was reported by Obeng et al. [28] and was generated using an inverted exon strategy (Figure 2). In this model, an exon 15 with the A>G substitution and a WT exon 16 were introduced in an inverted orientation within introns 16–17. Upon Cre induction, the inverted exons were recombined in the correct orientation in place of the endogenous WT exons through the use of different loxP sequences and, in theory, should be physiologically expressed from the endogenous locus. Although the *Sf3b1*^{K700E} allele was successfully inverted after Cre induction (with pI:pC treatment), it did not achieve heterozygous expression when the RNA levels were assessed. The authors presented results of RNA sequencing (RNA-Seq) of *lin*⁻*cKit*⁺ cells from the BM of three *Sf3b1*^{K700E/+} animals, where the mutant allele frequency was between 27% and 32% (see Figure S1A in Obeng et al. [28] and (see Supp Figure 1E-F in ref [23])). A similar level of subheterozygous mutant transcript expression was detected in a second recent report using this allele [24] and the phenotypes reported for these animals need to be considered with this caveat.

Monitoring of native hematopoiesis over 64 weeks of pI:pC-treated cohorts showed a progressive macrocytic anemia with decreased red blood cell counts from 20 weeks after activation of the K700E mutation. Corresponding with anemia, there was an increased level of plasma erythropoietin

in *Sf3b1*^{K700E/+} mice. This anemic phenotype was shown to be cell intrinsic through use of noncompetitive BM transplantation assays. There was no change in white blood cell or platelet counts. None of the *Sf3b1*^{K700E/+} mice was reported to develop MDS during the period of monitoring. At 64 weeks after pI:pC, analysis of the BM showed no change in cellularity, whereas there was a significant increase in the number of long-term HSCs (LKS⁺CD150⁺CD48⁻) and a decrease in the granulomonocytic progenitors (GMPs). In the spleen, there was an accumulation of erythroblast populations (Ter119⁺CD71^{high}) and a decrease in the more mature erythroid population (Ter119⁺CD71^{low}), indicating a terminal erythroid maturation defect. Interestingly, this effect was only reproduced in vivo in young mice (11 weeks after pI:pC) under stress (drug-induced hemolysis) or in vitro when BM progenitor cells (24 weeks after pI:pC) were treated with cytokines to induce erythroid differentiation. Morphological examination of the spleen showed an increased number of erythroid precursors and erythroid dysplasia. However, ring sideroblasts or elevated levels of iron deposition were not observed. There was a severe impairment (<20%) in the competitive repopulation capacity of *Sf3b1*^{K700E/+} cells (similar to Mupo et al. [29]), even though >95% chimerism was achieved in noncompetitive transplantation experiments.

Sf3b1^{K700E} “minigene” model (Mupo et al. [29]).

Mupo et al. [29] generated a conditionally activatable *Sf3b1*^{K700E/+} allele using a mini-gene approach (Figure 2). They engineered a loxP-flanked construct containing a splice acceptor site (from an Engrail-2 splice acceptor sequence), exons 12–15, intron 15, exons 15–19 (exons 12–19 were codon optimized sequence), intron 19, and the naturally occurring sequences for exons 20–25 including the 3'-untranslated region (UTR), followed by a SV40 polyadenylation signal (SV40 pA). This mini-gene was inserted between exons 11 and 12 of *Sf3b1* and the endogenous exon 15 downstream of the mini-gene was replaced with a mutated, synthetic exon 15 harboring an A>G mutation encoding K700E. Upon expression of Cre protein from *Mx1*-Cre induced by administration of pI:pC, the minigene was deleted, allowing the expression of the *Sf3b1* allele containing the mutated exon 15 (*Sf3b1*^{K700E/+}).

The allele was assessed in a heterozygous setting with one mutant allele and one WT allele, mirroring the patient setting. RNA-Seq analysis confirmed the 50% expression level of the mutated allele in whole BM and lineage-negative (*lin*⁻) cells from *Sf3b1*^{K700E/+} mice. Four weeks pI:pC treatment native hematopoiesis was assessed and a decrease in the percentage of phenotypic HSCs (LKS⁺CD34⁻Flk2⁻) was observed, but no change in the percentage of lymphoid-primed multipotent progenitors (L-MPPs), common myeloid progenitors (CMPs), GMPs, or myeloid-erythroid progenitors (MEPs). BM analysis indicated a mild myeloid bias

(Gr-1⁺Mac-1⁺) and impaired terminal erythroid differentiation (Ter119⁺CD71^{low}FSC^{low}). There was no overall survival difference between *Sf3b1*^{K700E/+} mice and WT littermates followed longitudinally for up to 83 weeks. Peripheral blood analysis revealed a progressive normocytic anemia with no change in white blood cell or platelet counts. None of the *Sf3b1*^{K700E/+} mice was reported to develop MDS during monitoring. Morphological analysis of the BM did not find evidence of either dysplasia or ringed sideroblasts (a feature of human *SF3B1* mutant MDS), despite increased iron deposits in BM macrophages. Peripheral blood morphology was not reported. Competitive BM transplantation was performed using both young (2 months) and old (12 months) recipients to determine the potential influence of aging recipients on donor cell engraftment. In both groups, *Sf3b1*^{K700E/+} cells showed poor chimerism, indicating impaired engraftment/repopulating potential. There was no difference between the young and old recipient groups with neither developing overt symptoms of MDS.

SRSF2. Serine-arginine rich factor 2 (SRSF2) binds to exonic splicing enhancers within the exonic sequence located near the 3' splice site and mediates exon inclusion and exclusion (Figure 1). Unlike *U2AF1* or *SF3B1*, *SRSF2* mutations occur near exclusively in myeloid malignancies and at or encompassing proline 95 (P95), with the most frequent mutation reported a proline to histidine substitution (P95H) [10,12,17]. *SRSF2* mutations occur in approximately 12–15% of MDS [9] and 28–52% of CMML [45–47] cases. Mutations are associated with poor outcomes, a shortened time to leukemic conversion, and a myelo-monocytic bias [48–50]. Three independently generated models of *Srsf2*^{P95H} have now been reported and these will be discussed in greater detail than those of *U2af1* and *Sf3b1* given our own work on this gene.

Srsf2^{P95H} knock-in models

***Srsf2*^{P95H} “gene duplication” model (Kim et al. [31]).** Kim et al. [31] reported the first *Srsf2*^{P95H} conditional knock-in model. The allele described approximated a gene duplication strategy (Figure 2). The targeting construct inserted a loxP site 87 base pairs (bp) upstream of the 5'-UTR sequence of the endogenous exon 1. The sequence containing exon 3 (3'-UTR) and an introduced SV40 polyadenylation sequence (SV40pA) were placed immediately after the endogenous exon 2 to generate an *Srsf2* allele that would express a WT protein. The second loxP was placed after the SV40pA. Downstream of the second loxP, the sequence of exon 1 containing the *Srsf2*P95H mutation (CG>AC) and the sequence for exon 2 were introduced into introns

2–3 (Figure 3). Very little data were presented describing the phenotypic effects of this *Srsf2*^{P95H/+} model on native hematopoiesis (i.e., not from transplantation recipients). The authors stated that none of the *Srsf2*^{P95H/+} mice developed AML up to 70 weeks of monitoring, although it is not certain if this refers to de novo-mutated animals or transplantation recipients.

The authors principally reported analysis from noncompetitive BM transplantation assays for the remaining studies. Subsequent studies with this *Srsf2* allele from the same authors have also used this approach [23,24]. At 18 weeks after pI:pC, the *Srsf2*^{P95H/+} recipients developed macrocytic anemia and leukopenia due to decreased B-cell numbers, whereas the platelet numbers remained normal. There was evidence of myeloid (hypolobated and hypogranulated neutrophils) and erythroid dysplasia in the peripheral blood/BM, but the estimated percentage of affected cells was not reported. Within the BM of the recipients, Kim et al. [31] reported that a major effect of the *Srsf2*^{P95H/+} mutation was an expansion of the HPC-2 phenotype (LKS⁺CD150⁺CD48⁺) within the hematopoietic stem and progenitor population. The frequencies of the HPC-2 population as reported by Kim et al. [31] in the control *Mx1-Cre Srsf2*^{+/+} animals was 0.0231%, significantly higher than that reported in the index reference for this population of 0.0032 ± 0.0014% in WT mice [51]. The frequency of this population may have been affected by both the BM transplantation model and pI:pC use. However, the ~10-fold difference in reported frequency was not reconciled and the representative flow cytometry gating strategy presented by Kim et al. [31] is significantly different from the gating strategy in the original report of this population [51]. The authors reported an increased frequency of cells in S-phase and early apoptosis within the BM. In a subsequent report from the same authors using this *Srsf2*^{P95H/+} allele and the *Mx1-Cre* model, the leukopenia and macrocytic anemia were largely absent (see Supp Fig 1E-F [23]). The inconsistency of the phenotype reported across the studies has not been addressed, but potentially indicates that the phenotype is not fully penetrant. Through analysis of RNA-Seq from both human patients and murine samples, the authors identified a group of mis-spliced candidate genes, with *EZH2* proposed as a key mis-spliced gene. *SRSF2*^{P95H} was proposed to lead to the inclusion of a premature stop codon in *EZH2*, resulting in a putative loss-of-function allele. This analysis posited a model that a single or small number of key mis-spliced genes may account for the phenotypes in the *SRSF2*^{P95H} mutant cells.

***Srsf2*^{P95H} “inverted exon” model (Kon et al. [32]).** Kon et al. [32] reported an *Srsf2*^{P95H} conditional knock-in model developed using an inverted exon strategy (termed FLEEx switch inversion technique; Figure 2). For the targeted allele, the 5'-UTR and exon

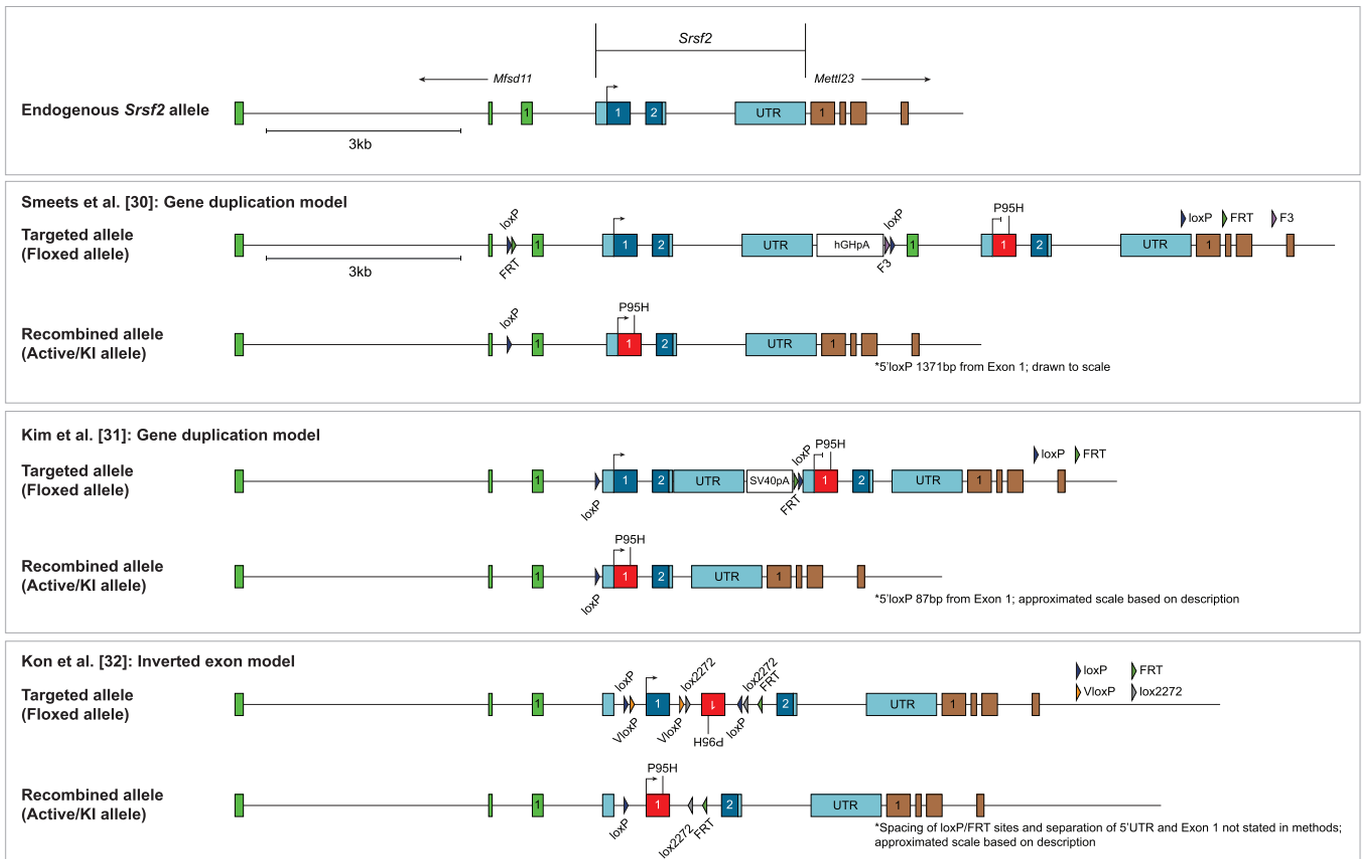


Figure 3. Detailed genomic architecture of the *Srsf2*(P95H) knock-in mutant models. (A) The endogenous *Srsf2* locus. Figure to scale provided. (B) The *Srsf2* allele described by Smeets et al. [30]. This allele is drawn to scale with the spacing of the inserted sequences as shown. (C) An approximation of the allele generated by Kim et al. [31] based on the description in the text. The 5'-loxP is 87 bp upstream of exon 1. The spacing and position of the remaining sequences is approximate. (D) An approximation of the allele generated by Kon et al. [32] using an inverted exon approach. The spacing and positions of the inserted sequences are approximations based on the description of the targeted allele. For (C) and (D), the modified locus depiction is approximate and based on the descriptions available in the studies reporting these alleles.

1 were separated. Exon 1 was flanked by loxP/VloxP sites at the 5' end and VloxP/lox2272 3' to exon 1. An inverted exon 1 containing the P95H mutation was inserted downstream of the VloxP/lox2272 in intro 1, followed by second set of loxP/lox2272 sites. After Cre activation, the inverted exon containing P95H “flips” into the correct orientation and the endogenous exon 1 is excised at the same time (Figure 3). RNA-Seq showed no expression from the modified allele in Cre-negative heterozygous animals, indicating that the floxed allele was likely null when not recombined. This was consistent with the failure to obtain animals that had the non-recombined P95H allele homozygosed even in a Cre-negative background [32]. Why this is the case is unclear at present. The targeting strategy itself may be the most significant contributor in this particular instance. The separation of the 5'-UTR from the first coding exon, in effect exonization of the UTR, and the numerous modifications around the coding exon and the sequence insertion within the first intron

may have had a significant effect on endogenous gene regulation.

The *Srsf2*^{P95H} conditional knock-in mice were crossed to *Vav1*-Cre mice to assess the effects of the heterozygous knock-in allele on hematopoiesis. Unlike the other described models, *Vav1*-Cre is a constitutive Cre and is active in the hematopoietic cells without the addition of an inducer such as pI:pC/IFN α (*Mx1*-Cre) or tamoxifen (CreER models). At approximately 15 weeks of age, the *Srsf2*^{P95H/+} mice developed macrocytic anemia, which remained stable. After observation up to 90 weeks of age, none of the *Srsf2*^{P95H/+} mice developed MDS or leukemia. A caveat with this model, as was described for the previous models describing inverted exon-based targeting methods, was that the mutant allele was expressed at 27–35% (mean 31%) based on RNA-Seq analysis of the *Vav1*-Cre *Srsf2*^{P95H} mice.

Upon analysis of the BM under native conditions, there was a reduction in the number of long-term repopulating

HSCs (LT-HSCs; LKS⁺CD150⁺CD48⁻), MPP (LKS⁺CD150⁻CD48⁻) and HPC-1 (LKS⁺CD150⁻CD48⁺) populations, which are associated with increased cell cycling. There was no change in the cellularity or lineage distribution of BM and spleen, with the exception of a mild B-cell differentiation defect in the BM. There was no evidence of dysplasia reported in the BM under native settings. The authors then performed noncompetitive and competitive BM transplantation assays of the *Srsf2*^{P95H/+} cells. There was engraftment in the noncompetitive transplantation setting, but the recipients of competitive BM transplantations had a significant reduction in chimerism when either whole BM or HSCs (LKS⁺CD34⁻) was used as the donor population. The impaired competitive transplantation potential is a common feature of all of the conditional splicing mutants to date. Extensive work demonstrated an engraftment defect of the *Srsf2*^{P95H/+} HSCs. The *Srsf2*^{P95H/+} recipients developed macrocytic anemia and leukopenia. In the BM, there was a significant reduction in LT-HSCs and most of the progenitor populations (MPPs, HPC-1, CMPs, CLPs). In addition, *Srsf2*^{P95H/+} recipients had a significantly higher proportion of cycling and apoptotic HSCs than controls. The lineage distribution was skewed toward myeloid at the expense of B-cell differentiation and dysplastic erythroid cells and dys-megakaryopoiesis were evident in the BM.

Analysis of RNA splicing using multiple purified hematopoietic populations from both native hematopoiesis and cells isolated from transplantation recipients was described. From these analyses, the authors could identify mis-splicing of a number of genes that had been identified in human *SRSF2* mutant patient samples, including *Csf3r*, *Gnas*, *Hnrnpa2b1*, and several novel differentially spliced genes such as *Atrx* and *Mllt10*, both of which are implicated in hematological malignancies. Despite using numerous approaches, the authors could not find evidence to support *Ezh2* as a differentially spliced candidate in the mouse. It was proposed that this may be due to species differences, with human *EZH2* having two CCNG motifs, whereas the mouse has a single CCNG motif [32].

***Srsf2*^{P95H} “gene duplication” model (Smeets et al. [30]).** Recently, we reported a third conditional *Srsf2*^{P95H} knock-in model [30] using a gene duplication method (Figure 2). The endogenous *Srsf2* locus including *Mfsd11* exon 1 and an introduced human growth hormone polyadenylation signal (hGHpA) was flanked by LoxP sites. The hGHpA was inserted downstream of the endogenous *Srsf2* 3'-UTR to prevent/reduce transcriptional read-through into the mutant allele in the absence of Cre. A duplication of the entire *Srsf2* locus and *Mfsd11* exon 1 was inserted downstream of the 3'-loxP sequence. The proline 95 (CCG) to histidine (CAT) mutation was introduced at the

duplicated *Srsf2* exon 1 (Figure 3). Upon Cre-mediated excision, the endogenous *Srsf2* locus was deleted and the duplicated locus containing the P95H mutation was retained. RNA-Seq confirmed the heterozygous expression at both the genomic and transcriptional level. Analysis of Cre-negative animals and animals not treated with tamoxifen (the CreER inducer used in these studies) demonstrated a low level of transcriptional read-through in the absence of activation of the P95H bearing allele. Long-term monitoring of Cre⁻ and non-tamoxifen-treated/recombined Cre⁺ animals demonstrated that this “leaky” expression was not pathogenic.

Multiple Cre lines were applied to understand the cell population in which P95H was required to be able to affect normal hematopoiesis: whole body/broadly expressed (*R26-CreER*^{T2} [52]), more specific to the HSC and primitive progenitor populations (*hScf/CreER*^{Tg/+} [53]), and a myeloid progenitor-targeted constitutive Cre line (*LysM-Cre*^{Ki/+} [54]). We analyzed the effect of *Srsf2*^{P95H/+} on native hematopoiesis at 20 weeks after activation of the P95H mutation. In all cases, Cre⁺ WT animals were used as controls and, for CreER models, these were tamoxifen-treated CreER⁺ *Srsf2* WT littermate controls. At 20 weeks after *Srsf2*^{P95H/+} activation, *Srsf2*^{P95H/+} mice developed macrocytic anemia and increased myeloid cells in the peripheral blood. The myeloid bias was more evident in the BM and was accompanied by compromised B lymphopoiesis. There was a reduction in erythropoiesis in the BM accompanied by increased splenic erythropoiesis. Within the phenotypic stem and progenitor populations, there was a reduction or trend to reduction in the number of stem cell populations using two different phenotypic methods (LKS⁺CD34/Flt3 [55–57] or LKS⁺CD105/CD150 [58,59]), whereas the more mature myeloid progenitor populations remained largely unchanged. The level of CD45RB, a previously characterized splicing target of *Srsf2* [21], was reduced in *Srsf2*^{P95H/+} splenocytes, indicating that altered RNA splicing was reflected in the proteome. A similar phenotype was seen in both *R26-CreER*^{T2} and *hScf/CreER*^{Tg/+} models, but not in the *LysM-Cre*^{Ki/+} model, indicating that the *Srsf2*^{P95H/+} mutation needed to arise within the primitive populations (including HSCs) to modify native hematopoiesis. RNA-Seq analysis of lin⁻cKit⁺eYFP⁺ cells (where enhanced yellow fluorescent protein [eYFP] marked cells and their daughters/progeny in which Cre was activated) isolated 20 weeks after activation of P95H demonstrated that the *Srsf2*^{P95H/+} mutation induced gene expression changes consistent with myeloid bias, loss of lymphoid potential, and transcriptional signatures found in MDS. We identified similarly mis-spliced transcripts as Kon et al. [32] and also could not define mis-splicing of *Ezh2* in the murine setting.

In agreement with the previous knock-in models of spliceosome mutations, the *Srsf2*^{P95H/+} cells exhibited a poor competitive engraftment potential. This appears to be a universal feature of the models described to date, with the work of Kon et al. [32] best characterizing this phenotype and demonstrating an engraftment defect in the mutant cells. This experimental result is confounding because *SRSF2* mutations are subjected to positive selection and are implicated in age-related clonal hematopoiesis [7,8,14]. However, we observed that the poor engraftment could be modified by altering the nature of the BM competitor used. Using the *hSc/CreER*^{Tg/+} model, we found that *Srsf2*^{P95H/+} cells could competitively engraft recipient animals and expand when transplanted with the age-/microenvironment-matched competitor cells (i.e., cells taken from the same BM but in which Cre was not activated based on R26-eYFP reporter marking of the cells).

Upon aging, nontransplanted *Srsf2*^{P95H/+} mice developed fatal MDS by ~12 months after activating the P95H mutation in both the R26-CreER^{T2} and *hSc/CreER*^{Tg/+} model. The disease in both models was highly comparable. When moribund, the mice presented with macrocytic anemia, myeloid bias (granulocytosis and monocytosis), and morphological dysplasia of myeloid and erythroid lineages in both the peripheral blood (>10–50% of cells) and BM, all characteristics of MDS/MPN. Analysis of a small cohort by exome capture demonstrated that there was a subclonal accumulation of mutations associated with human *SRSF2* mutant disease, including *Dnmt3a*, *Tet2*, *Phf6*, and *Ras* members, in the BM of the moribund animals [30,60]. Noncompetitive transplantation of the moribund BM recapitulated the MDS/MPN, but there was no evidence of progression to acute leukemia even with long-term monitoring of secondary recipients or with concurrent p53 deletion. Of the currently described *Srsf2* point mutant models, this is the only model to develop monocytosis and MDS/MPN in the setting of native hematopoiesis.

Discussion

Our understanding of the effects of spliceosome mutations comes from the detailed analysis of the mutational architecture of MDS, MDS/MPN, and related disorders in humans. In these settings, the mutation generally arises together with other mutations that combinatorially contribute to the disease manifestations and phenotypes [60]. In most cases, RNA-splicing mutations are not the sole causes of malignancies because other driver mutations are present and familial monogenic examples of RNA splicing mutations have not been described. However, isolated mutations in RNA splicing components can sometimes cause disease directly, such as in ~20% of *SF3B1*-mutant MDS in

which other driver mutations were not reported [17]. In the setting of CHIP, mutations in *SRSF2* and *SF3B1* have been identified with a range of variant allele burden, including VAFs of >10% [7,8,61]. These observations would suggest that, when present as the only driver mutation, spliceosomal mutations can confer advantages to the host cells. The development of murine models of human spliceosome mutations, both conventional knock-out models and humanized mutant models, has improved our understanding of how these mutations perturb normal hematopoiesis and ultimately promote malignancy. The establishment of tractable, autochthonous preclinical models has proven more challenging. The fidelity of the “humanized” murine models must be considered against the phenotypes associated with the same spliceosomal mutations in humans.

The independently generated “humanized” spliceosomal mutant mice described to date encompass a spectrum of phenotypes from very mild to overt MDS/MPN in the setting of native hematopoiesis (summarized in Table 1). The majority of models have also reported analysis of BM transplantation recipients. This is a useful strategy to limit the expression of the mutation to the transplantable hematopoietic cells, but is also a source of significant cellular stress. MDS, MDS/MPN, and related cancers arise in the absence of this extreme stress in the vast majority of humans. BM transplantation changes HSC clonal dynamics and contribution compared with native hematopoiesis and permanently damages and remodels the BM microenvironment itself [42,43,62–65]. The use of transplantation models also limits the capacity to understand the interplay of the mutant cells and a normal immune system, now appreciated to be an important aspect in cancer evolution and escape [66].

The models described demonstrate that the targeting strategy itself can have profound impacts on the utility of the model that is generated. The three models that have utilized an inverted exon approach have all observed ~30% variant allele frequency at the RNA level after locus recombination, demonstrating that, despite genomic heterozygosity, the expression from these modified alleles is not heterozygous. This caveat requires acknowledgement when these models are reflected against the human phenotypes associated with these mutations. The reasons that these alleles do not express heterozygously are not clear, but the results are consistent across three independent models and loci, indicating that this effect is most likely resultant of the targeting strategy itself. The inverted exon model has been successful in establishing murine models harboring the *Jak2*(V617F) mutation, in which the mice develop a myeloproliferative neoplasm resembling polycythemia vera, as occurs in humans with this

Table 1. Summary of the phenotypes of the “humanized” spliceosomal mutant mice described

Gene	Model	rtTA/Cre	VAF	Native hematopoiesis			Non-competitive transplantation			Competitive Transplantation
				Peripheral Blood/Spleen	BM	Dysplasia/Disease	Peripheral Blood/Spleen	BM	Dysplasia/Disease	
<i>U2af1</i>	Tet-On (Shirai et al. [33])	<i>R26-M2rtTA</i>	NA ^a	Not described			Leukopenia; ↓B cells; ↓CMP%; ^b ↑LKS*% ↓monocytes	No dysplasia, MDS, or AML up to 500 days post Tx	N/A	
	Inversion (Fei et al. [34])	<i>UBC-CreER^{T2}</i>	~30%	MEFs only; not further characterized						
	Mini-gene (Fei et al. [34])	<i>Mx1-Cre</i>	~50%	Macrocytic anemia; leukopenia; ↓B cells	↓LKS ⁺ ; ↓LTHSCs	<1% in BM; 0% in PB; no MDS or AML	Similar to native hematopoiesis		↓↓Engraftment	
<i>Sf3b1</i>	Inversion (Obeng et al. [28])	<i>Mx1-Cre</i>	~30%	Macrocytic anemia; ↓RBCs; ↑plasma EPO; ↑erythroblasts in spleen	↑LTHSCs; ↓GMPs	Erythroid dysplasia in spleen; no ring sideroblasts; no MDS (64 weeks post pI:pC)	Similar to native hematopoiesis		↓↓Engraftment	
	Mini-gene (Mupo et al. [29])	<i>Mx1-Cre</i>	~50%	Normocytic anemia	↓HSC%; ↑granulocytes; ↓mature erythrocytes	No dysplasia/ring sideroblasts; no MDS (83 weeks post pI:pC)	N/A		↓↓Engraftment	
<i>Srsf2</i>	Duplication (Kim et al. [31])	<i>Mx1-Cre</i>	~50%	Not described			Macrocytic anemia; leukopenia; ↓B cells	↑LKS; ↑LKS in S phase; ↑HPC2; ↓B cells	Neutrophil and erythroid dysplasia in PB; no AML (70 weeks)	↓↓Engraftment; ↑LKS; ↑HPC-2
	Inversion (Kon et al. [32])	<i>Vav1-Cre</i>	~31%	Macrocytic anemia	↓LTHSCs; ↓HPC-1	No dysplasia; no MDS or AML (90 weeks)	Macrocytic anemia; leukopenia; ↑granulocytes; ↓B cells	↓LTHSCs; ↓MPPs; ↓HPC1; ↓CMPs; ↓CLPs	Dysplastic erythroid and ↑megakaryocytes in BM; no MDS or AML	↓↓Engraftment; defect in homing
	Duplication (Smeets et al. [30])	<i>R26-CreER^{T2}</i> ; <i>hScf-CreER^T</i> ^{Tg/+} ; <i>LysM-Cre</i>	~50%	Macrocytic anemia; ↑granulocytes; ↑monocytes	↓LTHSCs; ↑%granulocytes; ↓B cells; ↓erythroid cells	MDS/MPN by ~12 months; no AML; ↑and dysplastic myeloid cells (10–50% PB); monocytosis in PB/transplantable to secondary recipients	Macrocytic anemia; leukopenia; ↑%granulocytes and monocytes; ↓B cells; ↑% monocytes in spleen	↑%granulocytes; ↑%monocytes	No AML	↓↓Engraftment in <i>R26-CreER^{T2}</i> model; myeloid bias; engraftment and ↑mutant cells in <i>hScf-CreER^T</i> ^{Tg/+} model

^aThe allele frequency is doxycycline dose dependent.

^bThe immunophenotypic population listed in this table is defined as following: LKS⁺, lineage⁻/c-Kit⁺/Sca-1⁺; LKS⁻, lineage⁻/c-Kit⁺/Sca-1⁻; LT-HSC, LKS⁺/CD150⁺/CD48⁻; MPP, LKS⁺/CD150⁻/CD48⁻; HPC-1, LKS⁺/CD150⁻/CD48⁺; HPC-2, LKS⁺/CD150⁺/CD48⁺; HSC (Mupo et al. [29]), LKS⁺/CD34⁻/flk2⁻; pre-MegE, LKS⁻/CD41⁻/FcγR⁻/CD150⁺/CD105⁻; pre-CFUE, LKS⁻/CD41⁻/FcγR⁻/CD15⁻/CD105⁺. %=percentage of this population.

mutation [67]. Although speculative, for alleles that provide a proliferative/survival advantage or are strongly positively selected, the inverted exon model may be feasible because even subheterozygous expression may be sufficient to afford the cell with the advantage of the mutant allele.

Another consideration is how foreign elements such as loxP sites are placed within the endogenous gene structure. Although literature describing the systematic assessment of locus modifications is limited, some principals have emerged with the evolution of the gene targeting methods and from the more recent high-throughput targeting vector generation methods [68–72]. These include the placing of loxP elements relative to the promoter and intron/exon junctions (>200 bp is generally advised); avoiding disruption of evolutionarily conserved regulatory elements and CpG islands that may be important for normal gene regulation; being cognizant of the intron and exon sizes when introducing new elements with caution when there are short introns/exons because this can affect splicing of the gene; and avoidance of modification to neighboring genes if possible. These generalized observations may be relevant to the alleles described for *Srsf2*, in which there are only two coding exons and the mutation occurs in the first exon. There is a CpG island that encompasses the 5' promoter region and 5'-UTR/exon 1, intron and exon 2 of *Srsf2* (UCSC genome browser mm10; chr11:116,852,080–116,853,903). The *Srsf2* allele described by Kon et al. [32], for example, used an inverted exon approach and was found to be transcriptionally inactive/silent when non-recombined. This allele involved significant modifications to the gene structure, including separation of the 5'-UTR and exon 1 coding sequence and insertion of multiple recombination elements in the introns [32]. The model developed by Kim et al. [31] placed the 5'-loxP element 87 bp from the start of the exon 1-containing sequence and modified the intron spacing significantly in both the pre- and post-recombined configuration [31]. All of these modifications can potentially affect the expression of the locus.

In humans, it is thought that spliceosome mutations arise within the HSC compartment [6,73]. It is well known that the cell of origin can affect the phenotypes from experimental leukemia models [74,75], so the use of different Cre drivers that target distinct and/or overlapping cell populations may be important in the presentation of the phenotypes in these models. Two of the *Srsf2* and one of the *Sf3b1* alleles have been assessed using the constitutively active *Vav*-Cre [24,32,76]. *Vav*-Cre is active in the hematopoietic cells, including HSCs, with evidence for activity also in endothelial cells and some activity in the reproductive tissues [41,77,78]. Multiple groups have used the inducible *Mx1*-Cre, a broadly active Cre that is very

efficient at recombining loxP-flanked alleles in hematopoietic cells, including HSCs and other tissues in the adult animal such as the cells composing the BM microenvironment [41,79,80]. *Mx1*-Cre requires administration of pI:pC to induce an innate immune-mediated interferon response, a known stimulus of cycling of HSCs [81,82]. Whereas this can be used to assess native hematopoiesis, the spliceosome mutant models described to date using *Mx1*-Cre have predominantly reported phenotypic/functional analysis from recipients of BM transplantations, which were treated after recovery with pI:pC. This approach is useful to restrict Cre activity in non-hematopoietic tissues because only the transplanted/engrafted BM will express Cre [39,83]. Such a strategy is effective for understanding the hematopoietic intrinsic effect of the mutation, but it introduces the stress of BM transplantation and changes the hematopoietic dynamics and BM microenvironment as a result [42,43,62–65]. Coupled with the administration of pI:pC 4–8 weeks after transplantation, this elicits substantive stress on the entire hematopoietic system. An alternative is to use CreER-based models in which tamoxifen is required to be administered to activate Cre activity. We utilized both the *Rosa26*-CreER^{T2} and *hScl*-CreER^T lines to allow widespread and relatively HSC-restricted activation, respectively, of *Srsf2*^{P95H} [30]. The CreER systems and the administration of tamoxifen are not benign, with evidence that toxicity of the CreER itself and tamoxifen administered at high doses subcutaneously can have direct effects on hematopoiesis [84,85]. These effects can be reduced by oral administration of tamoxifen, shortening the duration of exposure to tamoxifen, or by the use of heterozygous CreER alleles. The contribution of the specific Cre drivers to the phenotypes reported are likely secondary to the impacts of the locus specific genetic modifications themselves. However, a direct comparison of the same splicing mutation with different Cre strains would resolve this (e.g., *Rosa26*-CreER vs. *Mx1*-Cre). In all situations, the parallel analysis of littermate Cre⁺ WT and Cre⁻ gene-modified animals is an essential control for these studies whether using *Mx1*-Cre or CreER systems.

The models of spliceosomal mutation described to date have demonstrated a range of phenotypes. Definitive reasons for this remain uncertain, with some interpretations citing a lack of conservation of splicing between species. Although the spliceosomal proteins themselves are highly evolutionarily conserved, this is not an insignificant consideration [16]. By virtue of the higher level of conservation of the exons between mouse and humans (89–94%), the exonic splice-enhancer sequences are more highly conserved [86,87]. Despite this, there is still uncertainty regarding the identity of disease relevant mis-spliced candidates in human

samples and murine models. For example, in humans, it was reported that missplicing of *EZH2* [31] and, more recently, *CASP8* [24,88] occurred in *SRSF2* mutant cells. For *Ezh2* at least, this observation was not confirmed in multiple murine *Srsf2*^{P95H/+} mutant models in which other mis-spliced candidates identified in human samples could be confirmed [30,32]. The phenotypic recapitulation and conserved disease evolution and progression that occurred in the *Srsf2*^{P95H/+} model that we described indicates that the mouse is able to reproduce the core aspects of the human phenotypes associated with the *SRSF2* mutation. However, the identification of the specific mis-spliced genes that cause disease development has remained elusive. There are several possibilities that could account for this. First, transcriptomic analysis has yet to be performed on sufficiently purified disease initiating cells in both human and mouse and the heterogeneity of the samples assessed to date masks identification of the relevant mis-spliced transcripts. Compounding this is the fact that the computational analysis of splicing has yet to reach a consensus and the different computational methods yield distinct results. An alternative possibility is that subtle changes across many genes collectively contribute to the phenotype, rather than a small number of dramatically mis-spliced candidates. In such a possibility, it will require highly purified samples and sufficient sequencing depth to confidently identify these. Another possibility is that missplicing occurs within common cellular pathways in both species, but that the underlying individual genes are more variable and not necessarily conserved. An example of species conserved transcriptional and cellular consequence despite an apparent divergence of the individual transcripts can be seen in the A-to-I RNA-editing field and in the phenotypes of *ADARI* mutation in human and mouse [89–91]. Unlike the exonic sequences bound by *SRSF2*, intronic sequences are more divergent between human and mouse, with 66.1% conservation of branchpoint sequences across mammals [92]. The coordinated recognition of the 5' and 3' splice sites, together with branch-point and polypyrimidine tract selection, are critical to splicing fidelity. The utilization of highly conserved branch points is likely to be conserved to a greater extent across species than those with weaker recognition motifs, less conservation, or alternative branch-point spacing. These differences in the intronic sequences and motifs between species may contribute to the phenotypic differences between mouse and humans, in particular for *SF3B1*, which binds the branch point within intronic sequence of the pre-mRNA.

Murine models of human disease-associated mutations allow us to understand how these mutations individually contribute to disease initiation and maintenance. To provide this understanding, they must recapitulate as closely as possible the nature of the

mutation, the expression of the mutant allele, and the cell of origin. A critical assessment of the genetic modification strategies applied demonstrates that there are significant differences in the approaches that have been utilized. How the different targeting strategies and experimental approaches for individual gene mutations contribute to the phenotypes observed should be taken into consideration, along with the degree to which the model presents phenotypes that are consistent with the human disease carrying the same mutation.

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Author contribution statement

JX, MS, and CW wrote the original draft; JX, MS, ST, MW, LP, CW wrote, reviewed, and edited the paper.

Conflict of interest disclosure

The authors declare no competing financial interests.

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