Mutant U2AF1-expressing cells are sensitive to pharmacological modulation of the spliceosome

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Somatic mutations in spliceosome genes are detectable in ~50% of patients with myelodysplastic syndromes (MDS). We hypothesize that cells harbouring spliceosome gene mutations have increased sensitivity to pharmacological perturbation of the spliceosome. We focus on mutant U2AF1 and utilize sudemycin compounds that modulate pre-mRNA splicing. We find that haematopoietic cells expressing mutant U2AF1(S34F), including primary patient cells, have an increased sensitivity to in vitro sudemycin treatment relative to controls. In vivo sudemycin treatment of U2AF1(S34F) transgenic mice alters splicing and reverts haematopoietic progenitor cell expansion induced by mutant U2AF1 expression. The splicing effects of sudemycin and U2AF1(S34F) can be cumulative in cells exposed to both perturbations—drug and mutation—compared with cells exposed to either alone. These cumulative effects may result in downstream phenotypic consequences in sudemycin-treated mutant cells. Taken together, these data suggest a potential for treating haematological cancers harbouring U2AF1 mutations with pre-mRNA splicing modulators like sudemycins.

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Myelodysplastic syndromes (MDS) are the most common adult myeloid malignancy with up to 40,000 new cases diagnosed each year in the United States. MDS are a heterogeneous group of clonal haematopoietic stem cell disorders characterized by peripheral blood cytopenias and progenitor expansion; approximately one-third of patients will transform to a secondary acute myeloid leukaemia (AML) that has a poor prognosis. The only curative therapy is bone marrow transplantation, which is often not an option because of patient comorbidities. New treatment approaches are greatly needed. At least half of all MDS patient bone marrow samples harbour a mutation in one of several spliceosome genes, highlighting a potential genetic vulnerability. In addition, spliceosome gene mutations often occur in the founding clones of MDS tumours, providing an attractive target for elimination of all tumour cells. Spliceosome gene mutations are mutually exclusive of each other in patients, implying either a redundancy in pathogenic function or that a cell cannot tolerate two spliceosome perturbations at once. This in mind, we hypothesized that cells harbouring a spliceosome gene mutation would have increased sensitivity to further perturbation of the spliceosome by splicing modulator drugs. To examine this, we utilized sudemycin compounds that bind the SF3B1 spliceosome protein and modulate pre-mRNA splicing. We used sudemycin D1 and D6, which are synthetic compounds that have been optimized by several rounds of medicinal chemistry for their potent in vivo antitumour activity.

We examined the sensitivity of spliceosome mutant cells to sudemycin treatment, focusing on mutations in the spliceosome gene U2AF1, which have been identified in 11% of MDS patients, utilizing the S34F missense mutation most commonly found in our studies. Mutant U2AF1(S34F) expression has been shown by our group and others to cause altered pre-mRNA splicing in a variety of cell types, as well as altered haematopoiesis and pre-mRNA splicing in mice. In this manuscript, we provide evidence that U2AF1(S34F)-expressing cells are sensitive to the splicing modulator drug sudemycin. Haematopoietic cells expressing mutant U2AF1 show reduced survival and altered cell cycle in response to sudemycin D6 in vitro. In vivo treatment of U2AF1(S34F) transgenic mice with sudemycin results in an attenuation of mutant U2AF1-induced haematopoietic progenitor cell expansion that is associated with increased cell death. In addition, unsupervised analysis of whole-transcriptome sequencing (RNA-seq) finds that sudemycin D6 perturbs RNA splicing in both mutant U2AF1(S34F)- and U2AF1(WT)-expressing bone marrow cells; however, sudemycin D6 treatment further modulates mutant U2AF1(S34F)-induced splicing changes to create cumulative effects on cells in vivo. The cumulative RNA-splicing effects of sudemycin and mutant U2AF1 may contribute to the downstream phenotypic consequences we observe in vivo.

Results
Sudemycin alters RNA splicing in primary human CD34+ cells.
We first examined the pre-mRNA splicing alterations induced by sudemycin D6 in primary human haematopoietic cells. We treated CD34+ haematopoietic progenitor cells isolated from human umbilical cord blood with 1,000 nM of sudemycin D6 or dimethylsulphoxide (DMSO) vehicle control for 6 h in vitro and performed whole-transcriptome (RNA-seq) analysis (n = 6 each, Supplementary Fig. 1). We identified robustly altered gene expression and pre-mRNA splicing patterns induced by sudemycin, as shown by unsupervised clustering of samples using expressed genes (Fig. 1a) and pre-mRNA splice junctions (Fig. 1b), respectively. Our analysis identified 1,030 differentially expressed genes (FDR < 5%, |log2FC| > 1) and 18,833 dysregulated splicing events (FDR < 5%, |delta per cent spliced in or out| > 10%, Supplementary Data 1 and 2, respectively) that discriminated between sudemycin D6-treated samples and controls. Sudemycin D6 treatment induced altered pre-mRNA splicing with a bias towards increased exon skipping and intron retention (Fig. 1b). However, there was no apparent bias in the sequence motif surrounding splice acceptor sites of cassette exons that were alternatively spliced (Fig. 1c), in contrast to previously observed biases in sequences surrounding alternatively spliced junctions induced by expression of mutant spliceosome proteins U2AF1, SF3B1 and SRSF2 (refs 16–25).

To determine whether particular pathways are enriched for splicing perturbations, we applied GOseq to 6,278 genes with junctions significantly altered by sudemycin D6 treatment (FDR < 5%, |log2FC| > 2). While pathway enrichment was minimal (enrichment scores < 2), GOseq analysis indicated that pathways involved in pre-mRNA splicing, RNA processing and transport, cell cycle, as well as ATPase and helicase activity were enriched in splice junctions altered by sudemycin D6 treatment (FDR < 10%; Supplementary Data 1). Genes with sudemycin-altered expression were enriched in pathways involved in receptor and signal transduction activities (FDR < 10%; Supplementary Data 4).

Mutant U2AF1 cells have increased sensitivity to sudemycin.
To examine the effects of sudemycin D6 on haematopoietic cells expressing mutant U2AF1, we generated K562 human erythroleukaemia and OCI-AML3 AML cell lines that have stably integrated doxycycline-inducible, FLAG-tagged U2AF1(S34F) or FLAG-tagged U2AF1(WT) to control for U2AF1 overexpression (Supplementary Fig. 2a,b for K562; Fig. 2c,d for OCI-AML3). Mutant U2AF1(S34F)-expressing K562 cells showed reduced survival and lower IC50 (P < 0.0001, extra sum-of-squares F-test) relative to uninduced mutant U2AF1(S34F) and U2AF1(WT)-expressing control cells (Fig. 2a). These effects were also observed in human OCI-AML3 cell lines expressing mutant U2AF1(S34F) compared with U2AF1(WT)-expressing cells and other control cells (P < 0.003, extra sum-of-squares F-test; Fig. 2b). Reduced survival of K562 cells in the presence of sudemycin D6 is associated with an altered cell cycle profile: U2AF1(S34F)-expressing K562 cells had a decrease of cells in the S-phase and an increase of cells in the sub-G0/G1 and G2/M phases (Fig. 2c). Furthermore, MDS or AML cells with U2AF1(S34F) mutations treated in vitro with sudemycin D1, a sudemycin compound very similar to D6, showed an increased sensitivity to sudemycin (reduced S-phase) relative to control MDS/AML cells without spliceosome gene mutations (Fig. 2d). In contrast, treatment of MDS/AML patient cells with the chemotherapeutic drug daunorubicin (not predicted to disrupt splicing) showed no specificity for mutant U2AF1(S34F) samples compared with controls (Supplementary Fig. 2e). In addition, human CD34+ cells expressing U2AF1(S34F) showed increased sensitivity to another splicing modulator drug (E7107) similar to sudemycin (Supplementary Fig. 2f).

Sudemycin reduces mutant U2AF1 progenitor expansion in vivo.
We next examined the effect of sudemycin treatment in vivo on mutant U2AF1(S34F)-induced phenotypes using our previously described U2AF1(S34F) transgenic mouse model. We induced U2AF1(S34F) or U2AF1(WT) transgenes for 7 days in the bone marrow cells of transplanted mice (to study haematopoietic cell-intrinsic effects) and treated mice concurrently with sudemycin D6 (50 mg kg−1 per day) or vehicle for 5 of...
those days; see schema (Fig. 3a). Sudemycin D6 treatment of transplanted mice showed an attenuation of the previously described\textsuperscript{19} mutant U2AF1(S34F)-induced haematopoietic progenitor cell expansion by colony-forming unit (CFU-C) assay (Fig. 3b) and by flow cytometry for lineage-, c-Kit\textsuperscript{+}, Sca1\textsuperscript{+} (KLS) cells (Fig. 3c) when compared with control U2AF1 mutant mice treated with vehicle and mice transplanted with U2AF1(WT)-expressing bone marrow. The attenuation of mutant U2AF1-induced progenitor expansion by sudemycin-treated mice is associated with increased Annexin V\textsuperscript{+} staining of KLS cells (Fig. 3d).

**Sudemycin and U2AF1 (S34F) splicing effects can be cumulative.** To investigate the potential genotype-specific effects of sudemycin treatment on splicing isoform expression, we performed whole-transcriptome sequencing (RNA-seq) on U2AF1(S34F)- and U2AF1(WT)-recipient mouse bulk bone marrow cells following \textit{in vivo} U2AF1 transgene induction and treatment with sudemycin D6 (50 mg kg\textsuperscript{-1} per day for 5 days) or vehicle (Supplementary Fig. 3a,b). RNA was harvested 18 h after the last drug treatment (similar to described above; schema shown in Fig. 3a). Sudemycin D6 treatment at this dose and schedule does not markedly skew the mature lineage distribution within bulk bone marrow of mutant or wild-type (WT) U2AF1 transgenic mice (Supplementary Fig. 3c).

Using an unsupervised approach, we observed that sudemycin D6 perturbs splicing in both mutant U2AF1(S34F) and U2AF1(WT)-expressing bone marrow cells (Supplementary Data 5–9); this is visualized by the segregation of samples according to genotype and treatment within a principal component analysis (PCA) of cassette exon (Fig. 4a) and retained intron (Supplementary Fig. 4a) splicing events. Furthermore, the splicing bias observed in human cells treated with sudemycin (described above) is recapitulated in U2AF1(WT) mouse cells: sudemycin D6 induces exon skipping more often than exon inclusion relative to vehicle (388 of 657 significant (FDR < 10\%\textsuperscript{-1}, |ΔΨ| > 1\%) events; $P < 2 \times 10^{-6}$, one-sided binomial test), as well as intron retention more often than removal (98 of 145 significant (FDR < 10\%,
[ΔΨ] > 1%) events; P < 1.4 × 10⁻⁵, one-sided binomial test). As in human CD34+ cells, the sudemycin-induced changes were not associated with an apparent sequence motif (Supplementary Fig. 4b); however, we did observe the previously reported increase in a T in the −3 position of the intronic 3’ splice acceptor site of exons more commonly skipped in mutant U2AF1(S34F) cells16–19 (Supplementary Fig. 4c). In addition, we defined ‘high-confidence’ sets of U2AF1(S34F) and sudemycin targets, and subsets of those had a high validation rate in orthogonal experimental (NanoString26) and statistical (edgeR27) platforms (Supplementary Information and Supplementary Data 10 and 11).

Next, we examined potential interactions between the drug and mutation within cells, focusing on exon skipping events. Along these lines, we observed that the exon skipping effects induced by sudemycin D6 (relative to vehicle) within U2AF1(WT)-expressing cells are highly correlated with the drug effects in U2AF1(S34F)-expressing cells ($R^2 = 0.8$, $P < 2.2 \times 10^{-16}$, F-test, events significant in both comparisons (FDR < 10%), Fig. 4b). The vast majority of these events are concordant (in the same direction of induced change with similar magnitude) across genotypes (slope of the regression line = 0.75), suggesting that sudemycin treatment results in similar splicing alterations in these targets in both mutant U2AF1 and WT cells (Fig. 4b). We further assessed drug/genotype interaction using a statistical linear model: of 32,529 dysregulated splicing events (across all event types), only 136 showed statistically significant evidence of interaction (that is, synergy or antagonism; DEXSeq, FDR < 10%). However, when sudemycin D6 and mutant U2AF1(S34F) dysregulate a...
junction in the same direction (for example, both increasing exon skipping), this results in a cumulative effect in a sudemycin-treated mutant cell that is greater than the effect induced by sudemycin treatment of WT cells (indicated by red and blue colour in Fig. 4b–d). As an example, U2AF1(S34F) expression induces increased exon skipping in a 4932438A13Rik splice junction compared with U2AF1(WT) in vehicle-treated cells \( (\Delta \Psi = 0.205) \). Sudemycin D6 also increases exon skipping of the same junction in U2AF1(WT) \( (\Delta \Psi = 0.188) \) and U2AF1(S34F) \( (\Delta \Psi = 0.204) \) cells (that is, sudemycin-induced exon skipping is independent of genotype; Fig. 4c). Therefore, as expected, the cumulative effect induced by both mutation and drug treatment \( (\Delta \Psi = 0.409) \) exceeds the individual effects of mutant U2AF1 expression or sudemycin treatment alone, ultimately resulting in different levels of exon skipping in sudemycin-treated cells expressing mutant U2AF1 versus WT cells (Fig. 4c). The cumulative effect of mutation and drug can also be observed in splicing events that result in increased exon inclusion (Fig. 4d). Together, these data signify that the effects of sudemycin and U2AF1(S34F) on splicing in the same cell can be cumulative—that is, greater than the separate effects of drug or mutant.

Figure 3 | Sudemycin D6 treatment attenuates mutant U2AF1-induced progenitor cell expansion in U2AF1(S34F) transgenic mice. (a) Schema of sudemycin treatment of transgenic mice in vivo. Doxycycline-inducible U2AF1(S34F) or U2AF1(WT) transgenic mouse bone marrow was transplanted into recipient mice. Intrajugular (IJ) catheters were placed for IV drug infusions, which were performed over 4 h for 5 days. (b) Haematopoietic progenitor CFU-C colony-forming assay and (c) flow cytometry for haematopoietic stem and progenitor (HSPC) cell surface markers (c-Kit +, lineage −, Sca1 +, KLS, right panel) on U2AF1(WT)- or (S34F) mutant-recipient mouse bone marrow following treatment is shown \( (n = 6 \text{ for both U2AF1(WT) conditions, } n = 7 \text{ for vehicle-treated U2AF1(S34F), } n = 11 \text{ for sudemycin D6-treated U2AF1(S34F)}) \). (d) Annexin V + KLS cells were quantified in the bone marrow of mice following in vivo sudemycin treatment as described above \( (n = 7–9) \). Data pooled from two independent experiments (b–d); statistics calculated using two-tailed t-tests for each comparison shown; mean values with s.d. shown.
Figure 4 | Sudemycin D6 treatment alters splicing and gene expression in mutant U2AF1-haematopoietic cells. RNA sequencing and transcriptome analysis was performed on RNA harvested from mouse bone marrow cells expressing mutant U2AF1(S34F) or U2AF1(WT) following treatment of mice with sudemycin D6 (50 mg kg$^{-1}$) or vehicle control for 5 days (n = 5 per genotype and treatment). (a) PCA of normalized expression of skipped cassette exon events. (b) Correlation between sudemycin-induced changes relative to vehicle treatment in cassette exons of U2AF1(WT) cells ($ΔΨ_{WT} = Ψ_{WT,Sud} - Ψ_{WT,Veh}$) (FDR < 10%), and of mutant U2AF1(S34F) cells ($ΔΨ_{MT} = Ψ_{MT,Sud} - Ψ_{MT,Veh}$) (FDR < 10%), $R^2 = 0.8; P < 2.2 \times 10^{-16}$, F-test). Dashed line indicates similar effects in U2AF1(WT) and U2AF1(S34F) cells ($ΔΨ_{MT} = ΔΨ_{Sud}$). Colour scale indicates cumulative splicing changes of U2AF1(S34F) expression with sudemycin treatment (that is, $ΔΨ_{MT} = Ψ_{MT,Sud} - Ψ_{MT,Veh}$), with red being a positive change, blue being a negative change, and white being no difference between sudemycin-treated U2AF1(S34F) and U2AF1(WT) cells. (c,d) Delta PSI ($ΔΨ$) for each condition on the horizontal axis relative to vehicle-treated U2AF1(WT) cells ($ΔΨ_{WT,Veh} = 0$) for events that are significantly dysregulated across all five comparisons (see Supplementary Methods), have concordant sudemycin-induced dysregulation across genotype (that is, $ΔΨ_{WT} = ΔΨ_{WT} = ΔΨ_{MT}$ have the same direction), and in which the sudemycin effect is exacerbated by mutation. The cumulative effect on cassette exons induced by both mutant U2AF1 and sudemycin D6 treatment relative to vehicle-treated WT cells, $ΔΨ_{cum}$, may result in increased exon skipping (positive values, c) or increased exon inclusion (negative values, d). (e) PCA of normalized expression of expressed genes. (f) Correlation between sudemycin-induced changes relative to vehicle treatment (expressed as log$_2$ fold changes, FDR < 10%) in U2AF1(WT) and U2AF1(S34F) cells ($R^2 = 0.8; P < 2.2 \times 10^{-16}$, F-test). Dashed line indicates log$_2$ fold changes induced by sudemycin are the same in U2AF1(WT) and U2AF1(S34F) cells. Colour scale indicates cumulative contribution of mutant U2AF1 expression to sudemycin-induced gene expression changes, that is, log$_2$ fold change of gene expression altered by sudemycin in mutant U2AF1(S34F) cells relative to U2AF1(WT) cells, with red being a positive change, blue being a negative change and white being no difference. U2AF1 mutant (MT), sudemycin D6 (Sud), U2AF1 WT, vehicle (Veh), principal component 1 (PC1), principal component 2 (PC2), principal component 3 (PC3), per cent spliced in (PSI, $Ψ$).
Sudemycin induces gene expression alterations. Sudemycin treatment in vivo also results in altered gene expression in mutant U2AF1 mouse bone marrow cells compared with U2AF1(WT)-expressing control cells (Supplementary Data 12). As with splicing alterations, mouse bone marrow samples segregate by genotype and by treatment in an unsupervised analysis of gene expression (Fig. 4e). As seen at the junction level, there is a high-degree of correlation between sudemycin-induced, gene-level effects (direction and magnitude relative to vehicle) across mutant and WT genotypes ($R^2 = 0.8, P < 2.2 \times 10^{-16}$, F-test, events significant (FDR < 10%) in both comparisons, Fig. 4f). We also observed cumulative effects of mutation and drug on gene expression (colour in Fig. 4f), which may result in downstream cellular pathway changes. Using GOSeq, we identified pathways that were most enriched for differentially expressed genes in mutant cells treated with sudemycin D6 relative to the other genotype and treatment groups (FDR < 10%). We found mutant cells treated with sudemycin D6 were enriched in biologic pathways related to immune and inflammatory responses, antigen processing and presentation, cytokine production, leukocyte differentiation, cell death and apoptotic processes, when compared with the other genotype and treatment groups (Supplementary Data 13).

Discussion

We provide evidence suggesting that U2AF1(S34F)-expressing cells are sensitive to the splicing modulator drug sudemycin. Haematopoietic cells expressing mutant U2AF1 have reduced survival and cell cycle changes following sudemycin D6 treatment in vitro. In vivo, sudemycin D6 is capable of attenuating mutant U2AF1-associated expansion of haematopoietic progenitors in transgenic mice that is associated with increased cell death. In addition, while the effects of sudemycin D6 treatment on splicing and gene expression can be independent of the effects of mutant U2AF1 expression, sudemycin D6 treatment can also modulate splicing changes induced by mutant U2AF1(S34F) to create cumulative effects on cells in vivo. Together, these data indicate that mutant U2AF1(S34F)-expressing cells may have a therapeutic vulnerability to splicing modulator drugs such as sudemycin.

Previous studies using splicing modulator drugs, as well as non-pharmacological methods to target splicing factors, have indicated that the spliceosome is a promising therapeutic target for many cancer cell types. Recent studies have suggested that cancers with spliceosome gene mutations may have an increased sensitivity to splicing modulator drugs like sudemycin; this study provides further evidence to support this hypothesis. Specifically, we show that mutant U2AF1 cells are sensitive to in vivo treatment with sudemycin. Using unbiased RNA-seq, we show that sudemycin D6 effects on mutant U2AF1-associated splicing changes, resulting in different levels of transcript isoforms in cells expressing mutant U2AF1 compared with WT following sudemycin treatment. Ultimately, cumulative changes in isoform expression could be the cause of the mutant U2AF1-specific responses to sudemycin treatment that we observe in vitro and in vivo. Identification of critical downstream targets of sudemycin in vivo will likely require examination of various cellular populations within the bone marrow (including haematopoietic stem and progenitor cells) and harvesting cells at more immediate time points following drug treatment—both limitations of the current study. Future studies will focus on generating mutant U2AF1-expressing leukaemias in mice to test the efficacy of sudemycin on fully transformed haematopoietic tumours, as has recently been reported using E7107 to treat Srsf2(P95H)-expressing leukaemias.

Our data and others highlight several possible mechanisms for the sensitivity of spliceosome mutant samples to splicing modulator therapy. Pathway analysis of differentially expressed genes induced by sudemycin revealed enrichment in inflammatory signalling pathways. This is consistent with the enrichment in biological pathways related to cytokine and immune signalling observed following in vivo treatment of Srsf2(P95H) mutant mice with the splicing modulator E7107 (ref. 33), raising the possibility that spliceosome mutant-specific phenotypes observed in mice following splicing modulator drug treatment may be driven by an altered inflammatory response in mutant cells. Whether the altered inflammatory response in mutant cells treated with splicing modulator drugs is a direct result of mutant-altered splicing or gene expression could be explored in future studies. Alternatively, it is also possible that the cumulative effect of sudemycin and mutant U2AF1 expression on pre-mRNA splicing may simply create a state of ‘spliceosome sickness’ in cells by exceeding a tolerable threshold of splicing perturbations and their downstream consequences. Along these lines, other splicing modulator drugs have been shown to cause increased intron retention, R loop formation, DNA damage and cell death. Spliceosome gene mutations and splicing modulator drugs may both induce these consequences in a cell, and the cumulative effect of the drug and mutation may create a toxic intracellular milieu.

Exploring the clinical utility of splicing modulator therapies in MDS patients with spliceosome mutations who have failed current therapies is warranted and currently being pursued (NCT02841540), as these patients have few treatment options. Whether excessive toxicity will occur in WT cells treated with newer splicing modulators is a major question. Initial phase I clinical trial studies using a splicing modulator, E7107, showed toxicities in some patients with solid tumours, including ocular toxicity (NCT00459823, NCT00499499). The mechanism for the toxicity is not known. Moving forward, mutant U2AF1 and other preclinical models of spliceosome mutations will be valuable in further testing the in vivo efficacy and toxicity of drugs that modulate splicing through various mechanisms. Collectively, our data suggest that a mutant splicing factor together with a splicing modulator drug like sudemycin may create a unique cellular toxicity that could be exploited for therapeutic purposes.

Methods

Isolation and drug treatment of CD34⁺ haematopoietic cells. Mononuclear cells from human umbilical cord blood were separated by Ficoll gradient centrifugation (each n value is a pooled set of four individual umbilical cord blood samples). CD34⁺ cells were enriched using automACS-positive selection (CD34 MicroBead Kit, Miltenyi Biotec) according to the manufacturer’s instructions to achieve >90% purity of CD34⁺ cells. Following isolation, CD34⁺ cells were cultured in X-Vivo media (Lonza) with cytokines (SCF, Flt3L, IL3 and TPO) overnight. For whole-transcriptome sequencing with sudemycin, either sudemycin D6 (1,000 nM) or the vehicle DMSO was added to cells for 6h, and cells were then harvested for RNA. Tissue acquisition was performed per protocol approved by the Washington University School of Medicine Institutional Review Board.

RNA-seq of CD34⁺ cells treated with sudemycin. RNA was isolated from human CD34⁺ cells using the mirNAeasy Kit (Qiagen), and removal of genomic DNA was performed via a Turbo DNA-Free Kit (Ambion). Ribosomal RNA was removed using Ribozero (Epiconcept), followed by cDNA preparation and generation of stranded libraries using the TruSeq Stranded Total RNA Sample Prep Kit (Illumina). Sequencing was performed on the HiSeq2500 platform (Illumina) to generate 2 x 125 bp paired-end reads. RNA-seq data were deposited in NCBI dbGAP (phs000159.v9).
RNA-seq analysis of human CD34+ cells treated with sudemycin. Analysis of (stranded) RNA-seq data generated from primary human CD34+ haematopoietic cells treated with sudemycin D6 was performed using genome modeller system36. Reads were aligned to the human genome (hg19/NCBI build 37) using TopHat2 version 2.0.8, with annotations provided by Ensembl38 version 67. All downstream bioinformatic and statistical analyses, including calculation of P values using Fisher’s exact test and simulation, were performed in R39 and python. The heatmap of gene expression was normalized expression values output by DESeq2 package (version 1.6.3 (ref. 40)) that was subsequently z-scored. DESeq2 was used to detect differentially expressed genes, and gene-set enrichment analysis was then performed by applying Goseq version 1.34.0 to the differentially expressed genes (FDR < 0.05, log[FC] > 1, DESeq2). Additional details of RNA-seq analysis of gene expression are further described in the Supplementary Methods.

Alternative splicing events were computed via rMATs42 (version 3.0.8). We generated the heatmap of splice junctions using ‘per cent spliced in’ (PSI or α) values for exon skipping and intron retention events. For downstream analyses, namely reported regulated events and visualization of sequence contexts below, P values for events passing the filter were re-adjusted for multiple hypothesis testing using the method of Benjamini and Hochberg43.

A positive αPSI value indicates an event that was spliced in more often in the vehicle-treated relative to the sudemycin-treated samples. Sequence contexts of splice sites in exon skipping and intron retention events were visualized using seqLogo version 1.3.2.1. Dysregulated events visualized were those with αPSI > 10% and having a post-filtering re-adjusted FDR < 5% from rMATs. Dysregulated splicing junctions were determined by DESeq2 version 1.1.2.2. Only expressed junctions were analysed, and gene-set enrichment analysis was performed using the DESeq2 dysregulated events (FDR < 5%, FC > 3, DESeq). All of these are described in more detail in the Supplementary Methods.

Creation of inducible U2AF1(S34F) and U2AF1(WT) cell lines. Doxycycline-inducible FLAG-tagged WT U2AF1 or FLAG-tagged mutant U2AF1(S34F) lentiviral expression plasmids were previously described4, and lentivirus was generated in 293T cells with the packaging plasmids pMD-G, pMD-Lg and REV. Concentrated virus (multiplicity of infection (MOI) of 3 for K562, MOI of 5 for OCI-AML3) was used to transduce K562 cells (ATCC, CCL243) or OCI-AML3 (DSMZ, ACC 582). Transduced cells (marked by green fluorescent protein) were isolated by flow cytometry cell sorting. Expression of mutant or WT U2AF1 was induced using the indicated concentrations of doxycycline hyclate (Sigma, St Louis, MO) in water.

Cell line counting and BrdU incorporation assays. K562 (or OCI-AML3) cells in culture were seeded at ~200,000 cells per ml with different concentrations of Sudemycin D6 (or with DMSO control) following 48 h of initial doxycycline treatment to induce mutant U2AF1(S34F) or U2AF1(WT) expression. Cells were counted on Days 0, 3 and 5 of drug treatment using flow cytometry counting particle. Cell proliferation was measured by the manufacturer’s recommendations (modipropid iodide (Möllpore) to exclude dead cells from counts. Day 5 data were graphed in Graphpad Prism (Graphpad Software Inc.) using a nonlinear regression best fit line of the log (drug) versus response for each genotype and treatment. Statistical differences between curves were examined by the extra sum-of-squares F-test function in Graphpad Prism. For K562 cells, 3-bromodeoxyuridine (BrdU) incorporation was performed using the BrdU Flow Kit (APC. #596919, BD Biosciences) as per the manufacturer’s instructions using a 45 min pulse of BrdU of cells in culture on Day 5 of drug treatment.

Culture of MDS/AML cells and EdU incorporation assay. Primary MDS and AML cells were cultured on irradiated (1,000 rads) congenic WT recipient mice (C57BL/6 × 129S4/Scr/Jae)F1 (CD45.2.23) were used for culture of MDS and AML cells. Donor mice were between 8 and 12 weeks of age, and recipient mice ranged from 6 to 12 weeks of age; donor and recipient mice were sex-matched (both sexes were used in experiments). Donor chimerism was confirmed ≥6 weeks post transplantation to ensure engraftment of transgenic bone marrow. Post-engraftment, intrajugular treatment to induce mutant U2AF1(S34F) or U2AF1(WT) transgene on the day of surgery, and drug loading parameters were those of experiments that were chosen to allow for treatment with sudemycin or vehicle control. Investigators were not blinded to the group allocation during the experiment or analysis. Animals were excluded from analysis if their catheters did not remain patent during the entire treatment period. Mice were euthanised for analysis the day following the last drug infusion of treatment. Mice were randomised to treatment for the statistical comparison between groups. All mouse procedures were performed according to the protocols approved by the Washington University Animal Studies Committee.

Mouse haematopoietic progenitor cell assay. Methycelllose progenitor cell assay were performed using Methocult GF M3434 (Stem Cell Technologies). Bulk bone marrow cells were obtained from experimental mice, and red blood cell lysis was performed using 0.1% of 4% of 1,000 bone marrow cells per 1 ml media; each sample was evaluated in duplicate. Progenitor colonies (defined as ≥40 cells per colony) were counted following 7 days culture at 37°C with 5% CO2.

Mouse haematopoietic cell flow cytometry. For flow cytometry, all antibodies are from eBioscience (unless indicated), and catalogue number provided (if available). For haematopoietic progenitor/stem cells, we used the following antibodies (volume of antibody used in 200 μl of fluorescein-activated cell sorting (FACS) buffer for staining is also indicated): CD45.1-APC (#17-0453, 3 μl), CD45.2-PE (#12-0454, 3 μl), B220 (#13-0452, 0.5 μl), CD90.2 (#13-0302, 0.5 μl), CD19 (#13-0451, 0.5 μl), CD11b (#13-0451, 0.5 μl), Ter119 (#13-3921, 0.5 μl) and CD45R1(#13-0411, 1 ml), streptavidin secondary-eFluor655NC, c-KIT-APC (#11-0455, 200,000 cells per ml with different concentrations of sudemycin D6 was performed using the genome modelling program SeqScape version 1.0.2. Dysregulated events visualized were those with αPSI > 10% and having a post-filtering re-adjusted FDR < 5% from rMATs. Dysregulated splicing junctions were determined by DESeq2 version 1.1.2.2. Only expressed junctions were analysed, and gene-set enrichment analysis was performed using the DESeq2 dysregulated events (FDR < 5%, FC > 3, DESeq). All of these are described in more detail in the Supplementary Methods.

RNA-seq of mouse bone marrow cells in vivo. RNA was isolated and prepared as described above for human cells, using the miRNeasy Kit (Qiagen) followed by removal of genomic DNA via a Turbo DNA-Free Kit (Ambion) and ribosomal RNA using Ribozero (Epicenter). The RNA preparation and generation of stranded libraries was performed using the TruSeq Stranded Total RNA Sample Prep Kit (Illumina). Sequencing was performed on the HiSeq2500 platform (Illumina) to generate 2 × 126 bp paired-end reads. The RNA-seq data have been deposited in NCBI’s Gene Expression Omnibus and are accessible through the GEO Series accession number GSE89834.

RNA-seq analysis of mouse bone marrow cells in vivo. Analysis of (stranded) RNA-seq data generated from U2AF1(S34F) and U2AF1(WT) murine cells treated with sudemycin D6 or vehicle was performed similarly to the human CD34+ cell analysis described above, with some differences, as follows. Alignment again utilized genome model system, and reads were aligned to the mouse genome (mm9/NCBI build 37). ‘Per cent spliced in’ (PSI or α) values were calculated for all four conditions (U2AF1(S34F), U2AF1(WT) X (sudemycin D6, vehicle)) using rMATs. PCA was performed independently on events annotated by rMATs as spliced using the same methods. For the statistical comparison between groups, all mouse procedures were performed according to the protocols approved by the Washington University Animal Studies Committee.

To quantify the simultaneous effect of treatment and genotype for each splicing event, we calculated the change in per cent spliced in values for the four pairwise comparisons in which the genotype (alternately, treatment) was the same and treatment pair, but in which genotype pair (alternately, treatment) was the other. We refer to the unchanged condition as the ‘context’ and to the two conditions that differ as ‘A’ and ‘B’ and denote the corresponding change in the per cent spliced in values as Δα_ContextA – α_ContextB. In addition, we defined the cumulative effect of mutant treatment cell relative to WT, vehicle-treated cell as Δα - α_Context. Each of the five comparisons described were evaluated using rMATs, and P values for these events were then adjusted for multiple hypothesis testing as described above. Scattered plots were then plotted of event-specific splicing events dysregulated by sudemycin in both U2AF1(WT) and U2AF1(S34F) contexts (α_Context < 1%, FDR < 10% in both comparisons). To test the hypothesis that splicing differences between treatments (U2AF1(S34F), U2AF1(WT)) X (sudemycin D6, vehicle) are significant, we compared the distributions of fold changes in event splicing between treatments. For each of the five comparisons described were evaluated using rMATs, and P values for these events were then adjusted for multiple hypothesis testing as described above. Scattered plots were then plotted of event-specific splicing events dysregulated by sudemycin in both U2AF1(WT) and U2AF1(S34F) contexts (α_Context < 1%, FDR < 10% in both comparisons). To test the hypothesis that splicing differences between treatments (U2AF1(S34F), U2AF1(WT)) X (sudemycin D6, vehicle) are significant, we compared the distributions of fold changes in event splicing between treatments.
Non-additive (that is, super-additive synergetic or sub-additive antagonistic) interactions between drug and mutation were assessed in DEXSeq by comparing a generalized linear model that included main effects for drug and mutation with a second model that additionally included an interaction term representing drug/mutation synergy or antagonism. As above, splice site sequence contexts of exon-skipping events dysregulated by sudemycin D6 (\(\text{AP}^\text{WT}_{\text{1a}}\) > 1%, FDR < 10%) and corresponding unperturbed control events (\(\text{AP}^\text{WT}_{\text{1a}}\) < 0.1%, FDR > 50%) were plotted using sqLogo. Similarly, sequence contexts were displayed for events dysregulated (\(\text{AP}^\text{MT}_{\text{1a}}\) > 1%, FDR < 10%) or unaffected (\(\text{AP}^\text{MT}_{\text{1a}}\) > 1%, FDR > 50%) by U2AF1(S34F).

At the gene-level, PCA was performed as described above for human expressed genes. Genes differentially expressed within the above-described five conditions were determined via DESeq2. Scatterplots were then made comparing log fold changes of genes dysregulated by sudemycin in both U2AF1(WT) and U2AF1(S34F) contexts (FDR < 10% in both comparisons). Pathways enriched (FDR < 10%) for genes differentially expressed between U2AF1(S34F), sudemycin-treated cells and U2AF1(WT) and/or vehicle-treated cells were independently determined in a pairwise manner using GOseq. Again, expanded details of this approach can be found in the Supplementary Methods.

Data availability. All relevant data generated in this study are available at data-deposition sites. Human CD34+ cells treated with sudemycin D6 in vitro, data are available at NCBI dbGAP (phs000159.v9). For transgenic mice expressing mutant U2AF1(S34F) or U2AF1(WT) and treated with sudemycin D6 in vivo, data are available at Gene Expression Omnibus (GSE89934).

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Author contributions
The study was designed by: C.L.S., B.S.W., T.A.G. and M.J.W. Primary cell/cell line experiments performed by: C.L.S., M.T., R.T., S.K., J.S., A.C. and B.S. Mouse model experiments by: C.L.S., R.T., J.N.L., M.N. and S.K. RNA sequencing by: R.S.F., C.F. and M.O. Bioinformatics analysis by: B.S.W. Sudemycin drug development and synthesis by: C.L. and T.R.W. The manuscript was written and edited by: C.L.S., B.S.W., T.A.G. and M.J.W. All co-authors reviewed and approved the submission.

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