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Cancer-associated mutations in the spliceosome gene SF3B1 create a neomorphic protein that produces aberrant mRNA splicing in hundreds of genes, but the ensuing biologic and therapeutic consequences of this missplicing are not well understood. Here we have provided evidence that aberrant splicing by mutant SF3B1 altered the transcriptome, proteome, and metabolome of human cells, leading to missplicing-associated downregulation of metabolic genes, decreased mitochondrial respiration, and suppression of the serine synthesis pathway. We also found that mutant SF3B1 induces vulnerability to deprivation of the nonessential amino acid serine, which was mediated by missplicing-associated downregulation of the serine synthesis pathway enzyme PHGDH. This vulnerability was manifest both in vitro and in vivo, as dietary restriction of serine and glycine in mice was able to inhibit the growth of SF3B1MUT xenografts. These findings describe a role for SF3B1 mutations in altered energy metabolism, and they offer a new therapeutic strategy against SF3B1MUT cancers.

Introduction

Since their identification in 2011, mutations in spliceosome genes have been found in numerous human malignancies (1). Among the most broadly implicated genes is SF3B1, recurrently mutated in myelodysplastic syndrome (MDS), chronic lymphocytic leukemia, uveal melanoma, breast carcinoma, pancreas adenocarcinoma, lung adenocarcinoma, prostate adenocarcinoma, and many other cancers (2–8). Part of the U2snRNP complex, SF3B1 normally serves in the proper selection of 3’ acceptor sequences in pre-mRNA splicing reactions. In cancer, SF3B1 mutations create a neomorphic spliceosome protein that induces cryptic 3’ splice site selection by favoring an alternative branch point sequence (9, 10). The resulting aberrant splice junctions frequently lead to nonsense-mediated decay (NMD) and downregulation of mRNA from hundreds of affected genes (9). Yet, less is known about how these splicing alterations affect the downstream cellular processes and oncogenic properties of SF3B1MUT cells.

Despite this still nascent understanding of its biology, mutant SF3B1 has already become a target for novel therapeutic strategies, as many SF3B1MUT cancers have few effective treatments (11). However, unlike canonical oncogene targets like BCR-ABL, SF3B1 mutations do not seem to create “oncogene addiction,” as cancer cells with native SF3B1 mutation do not require the continued presence of mutant protein for their in vitro viability (11, 12). Therefore, rather than directly inhibiting the mutant protein, successful therapeutic strategies may come from a synthetic lethal approach, whereby the treatment targets a process whose disruption would be tolerated by normal cells, but detrimental to those with an SF3B1 mutation. Among the most attractive processes to target in this way are those that are nonessential to normal cells, as this will minimize toxicity of the treatment and facilitate its combination with other therapies. To date, processes whose disruption is deleterious to SF3B1MUT cells but nonessential to normal cells have not been identified.

Here we report investigations into the downstream functional consequences of SF3B1 mutations in human cells, providing...
evidence that mutant SF3B1 induces a specific remodeling of the proteome, cryptic splicing-induced downregulation of metabolic genes, and reprogramming of cellular metabolism. We also find that SF3B1 mutations create a heightened sensitivity to deprivation of the nonessential amino acid serine, suggesting a novel therapeutic strategy for targeting SF3B1\textsuperscript{MUT} cancers.

Results

Knockin of mutant SF3B1 alters the transcriptome of breast epithelial cells. SF3B1 is recurrently mutated in 2% of breast cancers and associates with poor prognosis in subgroups of patients, but the functional role of the mutation in this cellular context has been unexplored (13, 14). We therefore first aimed to characterize the isolated consequences of SF3B1 mutations in untransformed human breast epithelial cells. We used AAV-mediated gene targeting to engineer heterozygous knockins of the dominant K700E hotspot in spontaneously immortalized but nontumorigenic MCF-10A breast epithelial cells. To control for nonspecific effects of gene targeting, we also generated knockins of a synonymous base substitution at R702 that is a single nucleotide polymorphism (SNP) in the human population (15), and targeted WT (TWT) clones, where cells experience gene targeting, but remain WT (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI125022DS1). Four mutant and 4 control clones were independently isolated, and heterozygosity of the knockins was confirmed at both the DNA and mRNA levels (Supplemental Figure 1B).

Splice junction analysis of whole transcriptome RNA-seq from 2 mutant and 2 control clones revealed SF3B1\textsuperscript{K700E}-dependent alternative splicing. At a P value less than 0.01 and FDR less than 0.05, we identified 685 junctions whose usage was upregulated in SF3B1\textsuperscript{K700E} cells (Supplemental Table 1). Alternative 3’ splice site selection (A3SS) was the most frequent splicing event (49% of all junctions) (Figure 1A), with enrichment of cryptic acceptors 10–30 nucleotides upstream of canonical ones (Figure 1B). More than half (56%) of the SF3B1\textsuperscript{K700E}-enriched junctions were not annotated in transcriptome databases of normal human tissue (Figure 1A), compared with 12% of junctions overall, demonstrating enrichment of novel junctions in SF3B1\textsuperscript{K700E} cells. To orthogonally validate these RNA-seq data in all mutant and control MCF-10A clones, PCR was performed on candidate aberrant junctions in the proteome, cryptic splicing-induced downregulation of metabolic genes SF3B1\textsuperscript{K700E} knockins, a highly significant overlap (Figure 1E and Supplemental Table 1). We then compared the whole-gene mRNA expression of our model with that of 2 independent transcriptome data sets of human breast tumors: TCGA and METABRIC (16, 17). Of the genes downregulated in SF3B1\textsuperscript{K700E} MCF-10A cells, 32%, 21%, and 17% were also downregulated in SF3B1\textsuperscript{K700E} tumors from METABRIC, TCGA, or both data sets (Figure 1E and Supplemental Table 4), respectively. Notably, 66% (25/38) of genes downregulated and misspliced in MCF-10A cells were also among those downregulated in one or more of the SF3B1\textsuperscript{MUT} breast tumor data sets (Figure 1E and Supplemental Table 4), demonstrating the contextual robustness of missplicing-associated downregulation of mRNA. In contrast, overlap was lower between upregulated genes in SF3B1\textsuperscript{K700E} MCF-10A cells and breast tumors (Figure 1E and Supplemental Table 5). Taken together, these findings validate the disease relevance of our isogenic cell model and provide experimental evidence for determining which transcriptome alterations are directly caused by mutant SF3B1 in human breast tumors.

Mutant SF3B1 reshapes the proteome of breast epithelial cells. Although transcriptional effects of SF3B1 mutation have now been observed in several contexts, less is known about how these alterations affect the proteome. We therefore performed peptide mass spectrometry on isogenic MCF-10A cells using stable isotope labeling of amino acids in culture (SILAC). This experiment generated abundance values for 5,715 proteins. Using a log\textsuperscript{2} abundance ratio cutoff of ±0.5, we identified nearly twice as many proteins downregulated by SF3B1\textsuperscript{K700E} than upregulated: 381 versus 204 (Supplemental Tables 6 and 7). Integrating these results with our transcriptome analysis revealed that downregulated proteins exhibited missplicing of corresponding mRNA more frequently than upregulated ones: 13% (51/396) versus 3.0% (6/204), respectively ($\chi^2$ P < 0.01) (Figure 2A and Supplemental Figure 2A and Supplemental Table 1). This analysis further revealed a group of downregulated proteins with concomitant decrease in whole-gene mRNA (Figure 2B, red box) and another whose whole-gene mRNA was not significantly changed (Figure 2B, blue box). This suggests mutant SF3B1 may downregulate some proteins by direct suppression of mRNA (i.e., through missplicing-induced NMD) and others through translational or posttranslational mechanisms. Indeed, among downregulated and missspliced proteins, 27% (14/51) showed evidence of likely suppression by NMD, with reduction in whole-gene mRNA expression and aberrant junctions predicted to be NMD-sensitive (Figure 2, A and B, and Supplemental Table 6), and another 49% (25/51) had unchanged
Figure 1. Mutant SF3B1 alters the transcriptome of breast epithelial cells. (A) Classification of splice junctions upregulated in SF3B1K700E MCF-10A cells. A3SS, alternate 3′ splice site; A5SS, alternate 5′ splice site; EXI, exon inclusion; EXE, exon exclusion. Known refers to junctions annotated, and novel refers to those not annotated, in transcriptional databases of normal human tissue. (B) Nucleotide position of cryptic 3′ splice sites, relative to canonical AG, for all aberrant junctions in SF3B1K700E MCF-10A cells. (C) PCR validation of missplicing of the MUT and DLST genes in all 4 SF3B1MUT MCF-10A clones. Data are representative of 2 independent experiments. (D) Number of aberrant junctions per gene plotted against whole-gene mRNA expression in isogenic MCF-10A cells. Green, at least one NMD-sensitive junction; blue, NMD-insensitive junctions. (E) Transcriptional overlap between SF3B1MUT human breast tumors in TCGA and METABRIC data sets and SF3B1K700E MCF-10A cells. Overlap of aberrant junctions (upper) and whole-gene mRNA expression (lower) are shown. The smaller circles contain those genes with both altered whole-gene expression and aberrant junctions in SF3B1K700E MCF-10A cells.
Figure 2. Mutant SF3B1 reshapes the proteome of breast epithelial cells. (A) Aberrant junctions per gene plotted against protein expression determined by SILAC in isogenic MCF-10A cells. Green indicates genes with downregulated whole-gene mRNA and NMD-sensitive junctions. (B) Integration of protein and whole-gene mRNA expression in isogenic MCF-10A cells. Misspliced genes are in orange. The red box shows a population of genes with downregulated mRNA and protein, and the blue box shows a population with downregulated protein, but not mRNA. (C) Western blot validation of protein downregulation for misspliced genes MUT and DLST in isogenic MCF-10A cells. Data are representative of 2 independent experiments. (D) Quantification of cryptic SLC3A2 peptide abundance by integration of ion counts from mass spectra in isogenic MCF-10A cells obtained by SILAC. (E) RNA-seq correlation of cryptic splicing of SLC3A2 in isogenic MCF-10A cells, showing insertion of A-P-I-S between exons 5 and 6. PSI of the cryptic junction is shown. (F) GO pathway enrichment analysis of proteins differentially expressed in SF3B1<sup>K700E</sup> MCF-10A cells.
whole-gene mRNA expression and aberrant junctions predicted to be NMD-insensitive (Figure 2, A and B, and Supplemental Table 6). For the few proteins upregulated and misspliced, whole-gene mRNA was not increased, suggesting translational or posttranslational mechanisms may be at work (Figure 2, A and B, and Supplemental Table 7). Importantly, we orthogonally validated candidate protein alterations with Western blotting in both our MCF-10A and hTERT-IMEC isogenic models (Figure 2C and Supplemental Figure 2B). These data indicate that the predominant association of missplicing by SF3B1MUT with down, rather than up, regulation of misspliced genes observed at the transcript level also extends to the proteome in breast epithelial cells.

Because SILAC quantifies the most abundant fraction of the proteome—and because the splicing pattern of most genes is a single dominant isoform and several less abundant minor ones—the above analysis in fact reflects how mutant SF3B1 affects the expression of dominant protein isoforms (18, 19). We therefore next sought to determine whether any known or novel alternative splice junctions upregulated by mutant SF3B1 produce corresponding junction-spanning peptides. Known alternative splice isoforms of any kind could only be delineated in 2.5% (141/5,715) of our identified proteins, and this set of isoforms did not happen to contain any of the known junctions upregulated by mutant SF3B1. We did, however, find one peptide that spanned a novel cryptic 3′ splice acceptor site in the heavy chain of the integrin protein CD98, also known as SLC3A2. In each SF3B1K700E clone, abundance values of this cryptic peptide were 16-fold higher than the low-level expression in control cells (Figure 2D). The protein change was an insertion of 4 amino acids (A-P-I-S), and RNA-seq data showed the corresponding cryptic transcript resulting from insertion of the 12-bp intronic sequence preceding exon 5 that encodes these residues (Figure 2E). Mirroring the peptide-level quantification, the percent spliced in (PSI) of this cryptic splice junction was 34-fold higher (11 vs. 0.3%, P value = 5.89 × 10⁻⁷) in SF3B1K700E cells (Figure 2E). Thus, cryptic protein can be produced in breast epithelial cells following physiologic, heterozygous expression of mutant SF3B1.

Lastly, we subjected all protein expression changes, whether associated with direct missplicing or not, to Gene Ontology analysis. Among proteins downregulated in SF3B1K700E cells, there was a notable enrichment of pathways involved in energy metabolism, while upregulated proteins showed enrichment of pathways involving mRNA processing, mRNA splicing, and chromosomal localization of proteins (Figure 2F). Together, these data show that mutant SF3B1 reshapes the proteome in breast epithelial cells, producing missplicing-associated gene downregulation, cryptic protein, and altered expression of proteins in multiple cellular pathways.

Evidence for reprogramming of mitochondrial metabolism by SF3B1MUT in breast epithelial cells. Given the contextual robustness of missplicing-associated gene downregulation induced by mutant SF3B1, we next examined the metabolic pathways enriched in proteins downregulated in SF3B1MUT cells. One notably specific signal was in the ubiquinol-cytochrome c oxidoreductase complex, also known as mitochondrial complex III of the electron transport chain (ETC). It is formed by assembly factors UQCC1 and UQCC2, which stabilize mitochondrial cytochrome B (MT-CYB) and deliver it to other ubiquinol-cytochrome c reductase proteins to form mature complex III (Figure 3A) (20, 21). Among the 50 most downregulated proteins in our SILAC experiment, 7 were part of this complex (Figure 3A). None of these 7 had SF3B1MUT-dependent changes in whole-gene mRNA expression (Supplemental Table 2), suggesting their downregulation was at the protein level and not a direct result of missplicing. However, the assembly factor UQCC1 was heavily misspliced, with multiple aberrant transcripts containing cryptic terminal exons (Figure 3B). Though truncated, these aberrant transcripts would be NMD-insensitive, and their presence likely explains why the whole-gene mRNA expression (i.e., the sum of all isoforms) of UQCC1 was not lower by RNA-seq. Yet full-length, canonical UQCC1 transcript was indeed lower in SF3B1K700E cells (Figure 3B), consistent with a zero-sum competition model of splicing between U2 snRNP complexes with WT or mutant SF3B1 (9). UQCC1 was not a protein captured by SILAC, but Western blotting revealed it to be decreased in SF3B1K700E cells, along with other complex III proteins (Figure 3C). We then constitutively expressed canonical UQCC1, and this rescued not only UQCC1 levels but also levels of other complex III proteins in SF3B1K700E MCF-10A cells (Figure 3C). UQCC1 and UQCC2 were also decreased by SF3B1K700E in hTERT-IMEC cells, although the downregulation of mature complex III proteins was less pronounced, perhaps due to a greater baseline reserve of the UQCC1 assembly factor in this cell line (Supplemental Figure 3). SF3B1MUT breast cancers from TCGA also did not show reduced whole-gene UQCC1 expression by RNA-seq, but they did by microarray analysis, whose probes capture canonical, but not cryptic, mRNA (Supplemental Figure 4). UQCC1 was also the single most downregulated gene in SF3B1MUT tumors in the METABRIC data set, which used a microarray platform for mRNA analysis (Supplemental Figure 4). These data show that physiologic expression of mutant SF3B1 is sufficient to downregulate mitochondrial complex III levels, likely through missplicing of UQCC1.

In transferring electrons from ubiquinol to cytochrome c, complex III fuels oxidative phosphorylation as well as the regeneration of electron acceptors for aspartate biosynthesis (22, 23). We therefore sought to determine whether the observed SF3B1MUT-induced protein changes are accompanied by metabolic alterations. We first performed targeted metabolomics using liquid chromatography electrospray ionization tandem mass spectrometry (LC-ESI-MS/MS) in our isogenic cells, quantifying energy-relevant amino acids and metabolites involved in glycolysis and the TCA cycle. We validated the metabolomics platform using cells exposed to the biguanide metformin, which induced several known metabolic consequences of this drug in control MCF-10A cells, including increased lactate, α-ketoglutarate, and serine, along with decreased citrate, succinate, and aspartate (24). Interestingly, mutant SF3B1 did induce multiple alterations in energy metabolites, including decreases in the TCA cycle metabolites succinate, fumarate, and malate; increases in glucose and 3-hydroxybutyrate; and decreases in amino acids such as serine, glycine, and aspartate (Figure 3D). For orthogonal validation, we confirmed decreased aspartate in an independent SF3B1K700E MCF-10A clone using a fluorometric assay (Supplemental Figure 5). While attributions of metabolite abundance to particular protein expression levels are made difficult by the complex interplay of metabolite and enzyme dynamics (25), the finding of decreased aspartate in SF3B1MUT cells may nonetheless be suggestive of ETC impairment, as aspartate biosynthesis is
an essential function of mitochondrial respiration in multiple cell contexts (22, 23, 26–28). To test mitochondrial respiration directly, we measured oxygen consumption rate in 2 mutant and 2 WT MCF-10A clones with a Seahorse XF Analyzer, and this demonstrated that both basal and maximal cellular respiration were decreased by SF3B1MUT (Figure 3E and Supplemental Figure 6). In contrast, extracellular acidification rate was unaffected (Supplemental Figure 6). Together, these results provide evidence that SF3B1 MUT induces a reprogramming of mitochondrial metabolism characterized by a decreased capacity for cellular respiration in breast epithelial cells.
Figure 4. Mutant SF3B1 suppresses the serine synthesis pathway in breast epithelial cells. (A) Serine synthesis pathway. 3PG, 3-phosphoglycerate; PHP, phosphohydroxy pyruvate; pSer, phosphoserine. (B) RNA-seq of isogenic MCF-10A cells showing cryptic splicing of PHGDH leading to NMD. (C) Validation of cryptic splicing (upper) of PHGDH in MCF-10A and hTERT-IMEC isogenic cells, as well as downregulation of PHGDH protein (lower). Data are representative of 2 independent experiments. (D) Quantification (left) and representative images (right) of growth of isogenic MCF-10A cells with and without serine and glycine (SG). Scale bars: 100 μm. (E) Quantification of short-term (top) and representative images of long-term (bottom) growth of isogenic hTERT-IMEC cells with or without SG. For both breast epithelial cell models, data are representative of 2 independent experiments, and 3 biologic replicates were used per condition. ****P < 0.0001 for t test comparing all mutant to all WT, but independent clones are also shown. (F) Serine synthesized from 13C-glucose (M+3) is reduced in SF3B1K700E hTERT-IMEC cells at baseline and upon SG starvation. Three biologic replicates per condition from one experiment are shown. **P < 0.01 for t test comparing the M+3 fraction between WT and mutant cells in each media condition.
Mutant SF3B1 suppresses the serine synthesis pathway in breast epithelial cells. Another metabolic gene whose expression was strikingly affected by SF3B1MUT was phosphoglycerate dehydrogenase (PHGDH), an enzyme that catalyzes the conversion of 3-phosphoglycerate to phosphohydroxypyruvate as the rate-limiting step in de novo synthesis of the nonessential amino acid serine (Figure 4A). SF3B1MUT knockin breast epithelial cells exhibited enrichment of 2 different aberrant splice junctions in PHGDH, which are predicted to decrease total PHGDH mRNA expression by creating a cryptic terminal exon and inducing NMD, respectively (Figure, B and C). Indeed, whole-gene mRNA expression was lower by RNA-seq in SF3B1MUT MCF-10A cells (Supplemental Table 2), PHGDH protein was among the most downregulated proteins captured by SILAC (Supplemental Table 6), and Western blotting confirmed decreased PHGDH protein in both SF3B1MUT MCF-10A and hTERT-IMEC cells (Figure 4C). Moreover, in the TCGA and METABRIC data sets, PHGDH was among the most downregulated genes (Supplemental Table 4) in SF3B1MUT tumors.

Previous studies have shown that deliberate suppression of PHGDH or other components of the serine synthesis pathway renders mammalian cells less capable of growth and survival without exogenous serine, or, as is usually done in such studies, serine and glycine (SG), since serine can be synthesized directly from glycine (29–31). We therefore asked whether SF3B1MUT cells, with suppressed PHGDH levels, might exhibit a dependence on exogenous SG. When cultured in media without SG (−SG media), WT MCF-10A cells grew more slowly than in complete media, but cells nonetheless remained viable and mitotic (Figure 4D). In contrast, SF3B1K700E cells showed minimal growth without exogenous serine and glycine (Figure 4D). For hTERT-IMECs, WT cells were naturally more capable of growing without SG than WT MCF-10As, yet SF3B1K700E hTERT-IMECs still showed minimal growth without SG in both short-term proliferation and long-term colony formation assays (Figure 4E). Of note, the baseline growth in full media of SF3B1K700E breast epithelial cell knockins was slower than that of control cells (Supplemental Figure 7), but SG starvation disproportionately suppressed growth from this lower baseline in SF3B1K700E cells (Figure 4D, D, and E). These data demonstrate that SF3B1MUT increases dependence on exogenous serine and glycine for growth in breast epithelial cells.

PHGDH downregulation and serine/glycine auxotrophy suggest that serine synthesis could be impaired by SF3B1MUT. Indeed, our metabolomic analysis of isogenic MCF-10A cells did show decreased total intracellular serine and glycine levels in SF3B1K700E cells (Figure 3D). However, while suppression of the serine synthesis pathway decreases total intracellular serine and glycine in certain cell types grown in regular media, in most cell types suppression of the serine synthesis pathway will not decrease total intracellular serine and glycine unless the exogenous supply of serine is restricted (30, 32, 33). In contrast, the fraction of serine synthesized de novo in cells, as revealed by isotope tracing of 13C-glucose into M+3 serine, is ubiquitously decreased by suppression of the serine synthesis pathway in different cell types (33–35). Therefore, to directly determine whether serine synthesis is decreased by SF3B1MUT, we performed isotope tracing of 13C-glucose in our isogenic hTERT-IMEC cells in both +SG and −SG conditions. The total serine was lower in SF3B1K700E hTERT-IMEC cells at baseline compared with SF3B1WT cells, with a fold change of 0.581 (Supplemental Figure 8 and Supplemental Table 9), similar to SF3B1MUT MCF-10A cells with a fold change of 0.497 (Supplemental Table 8). In media containing SG (+SG media), the percentage of M+3 serine synthesized from 13C-glucose in SF3B1WT hTERT-IMEC cells was 2.7% of the total serine, comparable to baseline serine synthesis rates in regular media of breast epithelial cells seen in other studies (Figure 4F, Supplemental Figure 8, and Supplemental Table 9) (36, 37). In contrast, the percentage of M+3 serine in SF3B1K700E hTERT-IMECs was 0.19% of the total serine in +SG media, equal to only 4% of the M+3 serine in SF3B1WT cells (Figure 4F, Supplemental Figure 8, and Supplemental Table 9). Upon starvation of serine and glycine, both the absolute total serine and absolute M+3 serine were decreased compared with +SG conditions in SF3B1WT cells, but the percentage of total serine made up by M+3 serine increased to 77%, consistent with previous work on serine isotope changes upon SG starvation (Figure 4F, Supplemental Figure 8, and Supplemental Table 9). In breast cancer, SF3B1 mutations occur predominantly in hormone receptor–positive tumors, so we utilized an SF3B1K700E knockin of ER-positive T47D breast cancer cells. As in untransformed breast epithelial cells, PHGDH mRNA was misspliced and its protein was downregulated by SF3B1K700E in T47D cells (Figure 5A). We then assayed the 3D growth of these isogenic cells in conditions with and without SG. Notably, SF3B1K700E T47D cells showed no significant growth deficit in regular media compared with SF3B1WT cells (Figure 5A), in contrast to the baseline slower growth exhibited by untransformed SF3B1MUT breast epithelial cells (Supplemental Figure 7). Additionally, the growth of WT T47D cells was not significantly impaired by SG starvation in these conditions, indicating adequate de novo serine synthesis capacity in these cells (Figure 5A). In contrast, the size of SF3B1K700E T47D spheroids was reduced by 30%–40% in −SG media in 2 independent knockin clones, as compared with those grown in +SG, demonstrating that SF3B1MUT induces vulnerability to serine and glycine starvation in fully transformed breast cancer cells (Figure 5A).

Given that mutation-targeting cancer treatments are beginning to transcend the traditional boundaries of tissue of origin in their applications (40), we next sought to determine whether SF3B1MUT cancer cells from other tissue types might exhibit this vulnerability. SF3B1 mutations are more common in hematologic malignancies than in solid tumors, and we found that PHGDH...
In contrast, SF3B1\textsuperscript{K700E} HNT34 cells and SF3B1\textsuperscript{K666N} MUTZ3 cells were nonviable after 4 days without SG, and both cell lines showed aberrant splicing and decreased protein levels of PHGDH (Figure 5C). We also confirmed deprivation of serine is principally responsible for the suppressive effects of SG starvation in these cells (Supplemental Figure 10). Interestingly, a few SF3B1\textsuperscript{WT} leukemia cell lines also had lower PHGDH expression (MV411, Mono-Mac-6, KG-1), and these cells did show greater growth suppression without SG than the other SF3B1\textsuperscript{WT} cell lines (Figure 5C). This suggests that downregulation of PHGDH by mechanisms other than missplicing by mutant SF3B1 may also predict for reliance on extracellular serine. Taken together, these data show that in both
knockin and naturally mutated cancer cells of multiple cell lineages, SF3B1 mutation is associated with downregulation of PHGDH and heightened sensitivity to SG starvation.

**PHGDH overexpression and PHP supplementation rescue growth without exogenous serine.** To determine whether PHGDH downregulation is a mediator of this vulnerability, we reexpressed canonical PHGDH in SF3B1K700E breast epithelial cells and SF3B1H662Q Nalm6 cells, creating 2 independently derived populations of PHGDH-overexpressing cells for each cell model, along with 2 independently derived populations of GFP- or vector-overexpressing controls (Figure 6 and Supplemental Figure 11). In contrast to GFP, PHGDH was able to rescue cell growth without SG in SF3B1K700E MCF-10A cells (Figure 6A). A similar rescue of growth without SG by PHGDH was also seen in both SF3B1K700E hTERT-IMEC cells and SF3B1H662Q Nalm6 cells (Figure 6, B and C). These data support the idea that impaired serine synthesis due to missplicing-associated downregulation of PHGDH contributes to the vulnerability to SG starvation of SF3B1MUT cells. Consistent with this, we also found that growth of SG-starved SF3B1MUT cells could be rescued by supplementation with phosphohydroxypyruvate (PHP), the reaction product of PHGDH that serves as the substrate for subsequent steps of the serine synthesis pathway (Figure 6D).
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Figure 7. Cancer cells with naturally acquired SF3B1 mutation show vulnerability to serine and glycine starvation in vivo. (A) HNT34 cells were subcutaneously injected into NSG mice given +SG or –SG diets, and tumor volume measurements were obtained starting at 5 weeks, with 10 biologic replicates in each group. (B) MUT3 cells were subcutaneously injected into NSG mice given +SG or –SG diets and tumors were excised from euthanized animals and measured at week 4, with 4 biologic replicates in each group. ***P < 0.001, **P < 0.01, *P < 0.05 for t tests between dietary groups.

Discussion

Here we report that hotspot mutations in the spliceosome protein SF3B1 cause a reshaping of the proteome, reprogramming of cellular metabolism, and vulnerability to deprivation of the non-essential amino acid serine. These findings have several novel implications. While previous work has elegantly demonstrated the mechanism by which mutant SF3B1 induces aberrant splicing, understanding of its consequences on the proteome and downstream cellular processes is limited (9, 10). We found that the SF3B1*MUT proteome has a dominant pattern of protein down-regulation in genes affected by aberrant splicing. For many, the likely mechanism is NMD-induced mRNA downregulation. For others, it is likely translational or posttranslational, which might occur, for example, through impaired stability of proteins harboring indels created by cryptic splicing. We also found evidence that protein with a novel in-frame peptide insertion was expressed and enriched in SF3B1*MUT cells, in the form of cryptic SLC3A2. We suspect this was detectable because SLC3A2 is a highly abundant protein, but this finding does suggest that other, less abundant cryptic proteins may also be present in SF3B1*MUT cells. Whether—or how—these various protein alterations contribute to the oncogenicity of SF3B1 mutations is unknown. Tumor suppressor haploinsufficiency has been suggested as one possible mechanism (9), and there were tumor suppressor proteins downregulated by SF3B1*MUT in our study, such as NF1 and FANCD2 (Supplemental Table 7). An additional possibility is that cryptic proteins enriched in SF3B1*MUT cells could serve a gain-of-function oncogenic role, as has been suggested for the Notch pathway gene DVL2 in CLL (45). At the same time, such cryptic proteins might have implications for immune recognition of SF3B1*MUT cancers. While most cryptic transcripts induced by mutant SF3B1 are also detectable—albeit at much lower levels—and SF3B1*MUT cells, some could be sufficiently specific to SF3B1*MUT cells so as to produce cryptic peptides that would be recognized as nonself by the host immune system. If so, immunotherapeutic strategies aimed at targeting these putative missplicing-associated neo-antigens may afford therapeutic benefit, a testable hypothesis that we are currently investigating.

Our findings also demonstrate that SF3B1 mutations cause a specific deregulation of metabolic genes and reprogramming of cell metabolism, characterized by decreases in mitochondrial complex III formation, mitochondrial respiration, and flux through the serine synthesis pathway. In breast cancer, SF3B1 mutations rarely occur in triple-negative cases, and it is notable that this receptor subtype is highly dependent on the serine synthesis pathway, in some cases exhibiting amplification of PHGDH (36, 37). While inverse association does not necessarily equal mutual exclusivity, it could be that one reason acquisition of SF3B1 mutations is disfavored in triple-negative breast tumors is due to their serine synthesis pathway–suppressing effects. It is also worth noting that a defining feature of MDS driven by SF3B1*MUT is the ring sideroblast, which represents a pathologic deposit of iron in the matrix of structurally abnormal mitochondria that is accompanied by aberrant heme metabolism (4, 46). As suppression of PHGDH has

heightened sensitivity to SG starvation in SF3B1*MUT cells might be exploited therapeutically in vivo. Vousden and colleagues have shown that mice fed an SG-free diet will achieve a 60% reduction in blood SG levels, can tolerate the dietary modification, and in certain contexts will exhibit slower tumor growth (43, 44). We therefore performed xenograft experiments with SF3B1*K666N HNT34 cells in mice fed SG-free or amino acid–replete diets. We first employed HNT34 cells both because of their native SF3B1 mutation and because additional in vitro experiments showed their growth could be suppressed by partial serine depletion, a state which reflects the effects of dietary restriction in vivo (Supplemental Figure 12). As shown in Figure 7A, HNT34-derived tumors in mice fed an SG-free diet grew more slowly than those of animals that received a control, SG-containing diet. To determine whether this in vivo inhibitory effect could be seen in other SF3B1*MUT cells, we also tested the growth of SF3B1*K666N MUT3 cells. As shown in Figure 7B, the growth of MUT3-derived tumors was also decreased in mice fed SG-free food. These data thus provide evidence that SF3B1-mutant cells can be therapeutically targeted in vivo by dietary restriction of serine and glycine.
recently been shown to directly disturb mitochondrial homeostasis and heme synthesis in endothelial cells, it is possible that PHGDH missplicing and downregulation could contribute to mitochondrial dysfunction and altered heme metabolism in SF3B1MUT MDS (32). Moreover, impairment of the electron transport chain can itself produce ring sideroblasts, as exemplified by the sideroblastic anemia caused by germline mutations in the ETC gene NDUFB11 (47). This raises the possibility that missplicing and downregulation of the ETC gene UQCC1 by SF3B1MUT could contribute to ring sideroblast formation in MDS. We believe future studies are warranted to investigate how the specific disease phenotypes of SF3B1MUT cancers are influenced by aberrant splicing of metabolic genes.

It is also tempting to speculate that, in addition to shaping the metabolic behavior of SF3B1MUT cells, reprogramming of energy metabolism could provide an advantage in the initiation and progression of SF3B1MUT cancers. Indeed, metabolic reprogramming is a veritable hallmark of cancer that can drive tumorigenesis through rewiring of nutrient utilization and changes in metabolite-driven gene regulation (48). However, we believe it remains fundamentally mysterious how SF3B1 mutations confer a cloning advantage to cells, and that this mystery was highlighted by our observations that knocking mutant SF3B1 into untransformed breast epithelial cells slowed rather than transformed their growth (Supplemental Figure 7). Whatever they are, the advantages of SF3B1 mutations are likely to be contingent upon the genetic coconspirators and epigenetic differentiation of the cell, and it will be interesting to explore, in additional models, what environmental and cell-intrinsic conditions unleash the oncogenic properties of SF3B1 mutations, and whether metabolic reprogramming is involved.

Whether cause or consequence of cancer, the altered metabolic properties we observed in SF3B1MUT cells offered a novel vulnerability to deprivation of the nonessential amino acid serine. Previous studies have shown anticancer activity for serine restriction, but less is known about the molecular determinants of cancer responsiveness to this treatment (24, 31, 43, 44). Maddocks et al. showed that p53 mutation sensitizes HCT116 colon cancer cells to serine deprivation by favoring the diversion of residual serine into nucleotide rather than glutathione synthesis, but this group also found that a murine p53-mutant pancreas cancer cell line was resistant to serine deprivation, demonstrating the context-dependence of p53 mutation on this phenotype (43, 44). We observed heightened sensitivity to serine starvation associated with SF3B1 mutation in multiple contexts, including 2 independent breast epithelial cell knockin models, a breast cancer knockin model, a leukemia knockin model, and naturally mutated leukemia cells. We also found evidence that the mechanism underlying this vulnerability involves missplicing and downregulation of the serine synthesis pathway enzyme PHGDH, as restored expression of this enzyme was able to rescue growth without SG in SF3B1K700E breast epithelial cell knockins and SF3B1MUT leukemia cell knockins. Given the diversity of genes affected by SF3B1MUT missplicing, it would not be surprising if altered activity of other metabolic proteins in addition to PHGDH contributes to this phenotype. Nonetheless, we describe here a novel form of synthetic lethality for SF3B1MUT cells, in which a predictable missplicing event is tolerated by itself but not in the context of additional metabolic stress. The proposed therapy also has a target—exogenous serine—that is nonessential to normal cells, as serine starvation appears well tolerated in mice (31, 43). This favorable toxicity profile may, in turn, allow dietary serine restriction to be combined with other treatments, as has been done in mice with biguanides and glutaminase inhibitors (24, 44, 50). It is also conceivable that in vivo administration of a serine-depleting enzyme could produce even greater serine deprivation and anticancer activity while still remaining tolerable to the host. Indeed, asparaginase is a backbone of curative therapy for acute lymphoblastic leukemia, and arginine-depleting enzymes have shown signs of anticancer activity in patients (51). Whether serine deprivation will be tolerable or effective in humans is unknown, but our findings provide a rationale for investigating this therapeutic strategy in patients with SF3B1-mutant cancers.

Methods

Cell lines. HEK293T cells were from ATCC and grown in DMEM/10% FBS. MCF-10A cells were from ATCC and grown in DMEM/F12 with 5% horse serum, 20 ng/mL EGF, 10 μg/mL insulin, 0.5 μg/mL hydrocortisone, and 0.1 μg/mL cholera toxin. Gene-targeted MCF-10A derivatives were grown in the same media, except PIK3CAEs54K knock-ins (used as a control for the experiment shown in Supplemental Figure 7), which were cultured without EGF. hTERT-IMEC cells were a gift from JW Shay (UT Southwestern Medical Center, Dallas, TX) and were cultured in MCF-10A media, except horse serum was substituted with 2% charcoal dextran stripped FBS (52). HNT34, MUTZ, and CMLTI cells were from DSMZ. Nalm6 WT and SF3B1H62Q cells were from Horizon discovery, as were T47D WT, SF3B1K700E, and SF3B1K700K clones. K562, MOLM13, MONOMAC6, MV411, TFI, and KASUMI cells were gifts from LP Gondek (Johns Hopkins University, Baltimore, MD). KG1 cells were a gift from RA Casero (Johns Hopkins University, Baltimore, MD). All leukemia cells were grown in RPMI/20% FBS, with supplements as follows: 10 ng/mL GM-CSF for MUTZ3, 2 ng/mL GM-CSF for TF1, and 10 μg/mL insulin for MONOMAC6. T47D cells were grown in RPMI/10% FBS. SF3B1 genotype was confirmed in all lines by Sanger sequencing of exons 13–24. STR cell line authentication and mycoplasma testing was done upon receipt and routinely thereafter, with last testing done April 2018.

Gene targeting. Gene targeting was carried out using recombinant AAV technology as previously described (53). Briefly, AAV targeting vectors were produced by ligating PCR-generated homology arms spanning intron 12 to intron 17 of the SF3B1 gene into a pSEPT-containing AAV plasmid backbone (Agilent), which confers genetic resistance through a promoterless IRES-NEO cassette. Site-directed mutagenesis was then performed to generate the K700E and R702R alterations (Supplemental Figure 1). Virus was prepared by transfecting HEK-293T cells with pHelper, pRC (Agilent) and respective SF3B1 AAV plasmids. Breast epithelial cells were infected, and single genomeline-resistant clones were isolated and assayed for integration of the targeting cassette with PCR. Successfully targeted clones were exposed to Cre-expressing adenovirus to remove the neomycin cassette. All knockins underwent Sanger sequencing of genomic DNA and cDNA to ensure presence and transcription of the intended mutation. Primer sequences used for homology arm amplification are shown in Supplemental Table 10.

Transcriptome analysis. Total RNA was harvested from cultured cells using RNeasy and Qiashredder kits (Qiagen). RNA-seq libraries were constructed using the Illumina TruSeq RNA Sample Preparation
Kit v3. Sequencing was performed on an Illumina HiScanSQ system to obtain a total of more than 40 million paired-end reads of 101 bp in length per replicate. Reads were aligned using TopHat (version 2.0.10) against the GRCh38/hg38 version of the human genome, and assembly was carried out with Cufflinks pipeline. Differential mRNA expression between mutant and control cells was analyzed with DESeq2 from read counts obtained using HTSeq (htseq-count) and GENCODE (release 26) gene annotations. Four mutant MCF10A replicates (2 from the K700E clone and 2 from K700E2) and 4 WT replicates (2 from R702R1 and 2 from TWT1) were compared. Splice junctions were identified by RSeQC (version 2.6.4) and quantified using DESeq2, and PSI values for junctions were determined by dividing the read counts for a given junction by all read counts that included either the donor or acceptor site. Junctions with coverage of at least 10 reads in mutant samples were included, and a PSI threshold of 5 or more was employed. Novel junctions were those not found in major transcriptome databases, including KnownGene, AceView, and Ensembl.

Splicing events were classified using exon-intron annotations from the GRCh38/hg38 reference assembly. Because it was shown to comprise a very minor fraction of SF3B1WT-associated missplicing events (9), intron retention was not included in the analysis. A junction was classified as NMD-sensitive if it produced a premature termination codon (PTC) more than 55 nucleotides upstream of the last exon-exon boundary (54). Transcriptome data, including raw fastq files, identified junctions, and assembled transcript annotations, were deposited in NCBI’s Gene Expression Omnibus (GEO) and are accessible through GEO Series accession number GSE134475 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE134475). For splice junction usage between TCGA breast tumors and isogenic MCF-10A cells, all splicing events identified as upregulated in SF3B1WT breast tumors by Darman et al. were included except intron retention. For whole-gene mRNA expression comparison between isogenic MCF-10A and TCGA/METABRIC data sets, the latter was obtained through cBioportal and those genes with altered expression in SF3B1WT vs SF3B1WT tumors at a false discovery rate value of less than 0.05 were included. The criteria for including SF3B1 mutations from the TCGA, METABRIC, and Beat AML data sets in the SF3B1WT group were that the mutations are recurrent in primary human cancers, create in-frame amino acid changes, and are located in the HEAT repeats domain, as DeBoever et al. previously showed that mutations outside the HEAT repeats do not exhibit the same cryptic splicing pattern (55). Validation of aberrant junctions was done with endpoint PCR on cDNA from cells using primers designed to amplify both the cryptic junction of interest and its paired canonical junction. These primers sequences are provided in Supplemental Table 10.

Proteome analysis. SILAC-based mass spectrometry of MCF-10A cells was performed as previously described (9). Briefly, 2 independent K700E clones and a TWT clone were propagated for more than 7 population doublings in DMEM/F12 SILAC media deficient in both l-lysine and l-arginine (Thermo Fisher Scientific) and supplemented with light lysine (K) and arginine (R) for light, ^13^H4-K and ^13^C6-R for medium state, and ^13^C6/N2-K and ^13^C6/N4-R for heavy state labeling (Cambridge Isotope Laboratories). Cells were lysed in 9M urea in ammonium bicarbonate buffer and equal amounts of proteome from the 3 SILAC-labeled states were mixed. Proteome was subjected to reduction and alkylation with 10 mM DTT and 20 mM iodoacetamide, respectively. Sequencing grade trypsin (Promega) was added in a 1:20 ratio followed by overnight trypsin digestion at 37°C. Peptides were prefractionated by basic RPLC chromatography using Waters X-treme C18 column, and 12 fractions were collected. Each fraction was subjected to LC-MS/MS analysis on a Thermo Orbitrap Fusion Lumos mass spectrometer. Peptides were separated on an EASY-Spray analytical column (2 μm, 50 cm, Thermo Scientific) over a 180-minute gradient. The data were acquired in a data-dependent manner in top speed mode over 3 seconds. Peptides were fragmented in higher-energy collisional dissociation (HCD) mode at 35% collision energy. Dynamic exclusion of 30 seconds was included in the method. Data were analyzed in 2 ways. First, for routing, protein quantitation spectral data were searched against Human RefSeq data set (release 73) using SILAC-based quantitation method. Second, for identification of novel splice junction peptides, the spectral data were searched against a 3 frame translated transcriptomic data obtained as a part of the same study. Peptide identification and quantification was done using Proteome Discoverer suite (version 2.1, Thermo Fisher Scientific). Peptide spectral matches qualifying 1% FDR by decoy database search method were accepted. Western blotting was performed as previously described (56). The following antibodies were used: anti-UQCC1 (Sigma-Aldrich, HPA034875), anti-UQCC2 (Sigma-Aldrich, HPA039111), anti-UQCRF1 (Abcam, ab14746), anti-UQCRCl1 (Abcam, ab110252), anti-UQCRQ (Abcam, ab110255), anti-MUT (Abcam, ab133672), anti-DLST (Abcam, ab187699), anti-PHGDH (Sigma-Aldrich, HPA021241), anti-β-Actin (Cell Signaling, 4970S), anti-GAPDH (Cell Signaling, 5174).

Gene ontology analysis. Proteins identified as downregulated or upregulated at log2 abundance ratios of ±0.5 in SF3B1WT MCF-10A cells by SILAC were submitted to GO biological process enrichment analysis through the Gene Ontology Consortium (http://geneontology.org) (57). Overexpression experiments. For UQCCI overexpression, canonical UQCCI was PCR amplified from cDNA made from MCF-10A cells and cloned into the lentiviral vector pLX301 (Addgene plasmid 25895). The pEXP7-TET cassette from the BP Clonase Enzyme Kit (Thermo Fisher Scientific) was cloned into pLX301 as a vector control. Lentivirus was produced by transfection of 293T cells with pMD2.6 (Addgene plasmid 12259), psPAX2 (Addgene plasmid 12260), and the pLX301 derivatives. For PHGDH overexpression in MCF-10A and hTERT-IMEC cells, we made lentivirus as above, but with plJM5-WT PHGDH lentiviral vector (Addgene plasmid 83901). For GFP control virus, we replaced the PHGDH ORF in plJM5-WT PHGDH with GFP from pmaxGFP (Lonza) and produced lentivirus otherwise as above. Breast epithelial cells were then infected with lentivirus, selected with puromycin (UQCCI) or hygromycin (PHGDH), and collected as independent polyclonal pools of stably expressing cells. For PHGDH overexpression in Nalm6 cells, the full-length PHGDH open reading frame was PCR amplified from cDNA from MCF-10A cells, inserted into the lentiviral vector pLX301, and lentivirus produced as above. The pEXP7-TET-pLX301 virus was used as a vector control, as above. Independent polyclonal Nalm6 pools of transduced cells were selected with puromycin.

Metabolite quantification. For MCF-10Ais, gene targeting control cells (R702R1) were grown over 4 days with or without 5 mM metformin, and SF3B1WT knockin cells were also grown for 4 days, but seeded at a higher starting density so that cell confluence was similar by the end of the experiment. Media was changed at 48 and 72 hours. At the time of harvest, cell counting was done on extra replicates of cells cultured in the same conditions and used just for the purpose of determining the volume of trypsinized cells needed to pellet 5 x 10^6
cells in the samples of interest, to minimize processing time. Cells were trypsinized for approximately 1 minute with 0.25% trypsin, and the predetermined volumes were spun and pellets snap-frozen in liquid nitrogen. The protein content of samples was then quantified using the remaining trypsinized cells with Pierce 660 per the manufacturer’s protocol for use in normalization of final metabolite levels. The frozen pellets were shipped to Biocrates for extraction and quantification of energy metabolites by mass spectrometry. Orthogonal validation of aspartate levels was performed using similar culture conditions, but with independent isogenic MCF-10A clones using the Aspartate Assay Kit from Sigma-Aldrich (MAK095) per the manufacturer’s protocol.

For hTERT-IMECs, WT and SF361 knockout cells were grown in 10-cm dishes in +SG and –SG media for 42 hours, and media was refreshed, but with 1°C-glucose for an additional 6 hours. Media was then withdrawn from the dishes and ice-cold 80% MeOH was added directly to cells. Cells were scraped off the dish and flash frozen in liquid nitrogen. Samples were vortexed and stored at –80°C for at least 2 hours. Media was then then withdrawn from the dishes and ice-cold 80% MeOH was added directly to cells. Cells were scraped off the dish and flash frozen in liquid nitrogen. Samples were vortexed and stored at –80°C for at least 2 hours to precipitate protein. Protein obtained from cells during metabolic extraction was quantified using BCA per the manufacturer’s protocol (MilliporeSigma) and protein levels were used for normalization of total ion abundance levels in the samples. Metabolite-containing supernatant was isolated after centrifugation at 15,000g for 10 minutes and dried under nitrogen gas and frozen at –80°C for subsequent analysis by liquid chromatography mass spectrometry (LC-MS). Dried metabolite extracts were resuspended in 50% acetonitrile solution. LC-MS-based metabolomics profiling was performed on an Agilent LC-MS system consisting of an Agilent 1290 Infinity Binary UHPLC pump and a 6230 time-of-flight mass spectrometer. Chromatographic separations were performed using an Agilent 1290 ultra-high-performance liquid chromatography system with a wellplate autosampler. An ion pairing method was developed using a C18 column (Agilent Zorbax Extend C18, 2.1 × 150 mm, 1.8 μm) with tributylamine as an ion-pairing agent, which enables the detection of most of the metabolites in central carbon metabolism. The LC parameters were as follows: autosampler temperature, 4°C; injection volume, 2 μL; column temperature, 40°C; and flow rate, 0.25 mL/min. The solvents and optimized gradient conditions for LC were as follows: Solvent A, 97% water/3% methanol containing 5 mM TBA and 5.5 mM acetic acid; Solvent B, methanol containing 5 mM TBA and 5.5 mM acetic acid; a nonlinear gradient from 0% B to 99% B in 22 minutes with 5 minutes of post-run time. A 6520 accurate-mass Q-TOF LC-MS system (Agilent) equipped with a dual electrospray (ESI) ion source was operated in negative-ion mode for metabolic profiling. The optimized ESI Q-TOF parameters for MS experiments were: ion polarity, negative; gas temperature, 325°C; drying gas, 10 L/min; nebulizer pressure, 45 psi; capillary voltage, 4000 V; fragmentor, 140 V; skimmer, 65 V; mass range, 50–1100 m/z; acquisition rate, 1.5 spectra/s; instrument state, extended dynamic range (1700 m/z, 2 GHz). Spectra were internally mass calibrated in real time by continuous infusion of a reference mass solution using an isotropic pump connected to a dual sprayer feeding into an electrospray ionization source. Data were acquired with MassHunter Acquisition software (Agilent Technologies).

For quantification of oxygen consumption rate, isogenic MCF-10A cells were seeded at 28,000 cells per well in Seahorse XF96 Analyzer plates in regular media and incubated at 37°C with 5% CO2 overnight. Media was then removed, cells were washed, Seahorse media (RPMI + 175 mm glucose + 1 mM pyruvate + 2 mM glutamine) was added to wells, and the plate was incubated at 37°C without CO2 for 1 hour. The plate was then loaded into the Seahorse machine and readings of oxygen and pH levels were taken before and after injections of 3 μM oligomycin, 4 μM FCCP, 2 μM antimycin, and 2 μM rotenone. Oxygen consumption rate and extracellular acidification rate were analyzed using Wave 2.4 software.

**Serine and glycine starvation**. Amino acid-free RPMI powder (US Biological, R9010-01) was individually reconstituted with RPMI-standard concentrations of amino acids (Sigma-Aldrich) as needed for experimental conditions. All serum used for starvation experiments was dialyzed FBS (dFBS, Thermo Fisher Scientific, 26400044). For MCF-10A and hTERT-IMEC cells, which were found capable of growing in RPMI, supplements for breast epithelial cell growth (EGF, insulin, cholera toxin, hydrocortisone) were added to reconstituted RPMI, with the same concentrations used for DMEM/F12 as detailed above, except 5% horse serum was substituted with 5% dFBS. MCF-10A isogenic cells were grown in +SG or –SG media for 9 days with media changes every 2 days, and viable cells were then counted with a Vi-Cell XR cell counter. Isogenic hTERT-IMEC cells were grown in +SG and –SG media for 6 days, at which point viable cells were counted. MCF-10A and hTERT-IMEC cells overexpressing either GFP or PHGDH were cultured in +SG or –SG in the same manner, as were phosphohydroxypropruvate (PHP) rescue experiments, but with increasing concentrations (0–1 mM) of PHP. For Nalm6 isogenic cells, RPMI + 20% dFBS with or without SG was used, cells were seeded at a starting density of 106 cells/mL, equal volumes of +SG or –SG media were added at 48 hours and 96 hours, and cell mass/viability was determined on day 6 with alamarBlue, per the manufacturer’s protocol. Other leukemia cell lines were cultured in +SG or –SG media using RPMI + 20% diazoyl FBS and, where required, the same supplements (GM-CSF, insulin) as in their routine culture media. Cells were seeded at a starting density of 106 cells/mL, an equal volume of additional +SG or –SG media was added at 48 hours and 96 hours, and cell mass/viability was quantified by alamarBlue on day 4. For starvation of serine-vs.-glycine-vs.-both experiments, HNT34 and MUTZ3 cells were seeded in respective media at 106 cells/mL and viable cells counted at 48 hours. For gradient starvation of serine, HNT34 cells were seeded at 106 cells/mL in media containing increasing increments of serine, from 0 to the standard RPMI concentration of 285 μM, and viable cells were counted at 96 hours. For isogenic T47Ds, cells were seeded into 96-well low attachment plates (Corning) at a density of 5000 cells per well in 200 μL of RPMI/10% dFBS with or without SG. Plates were spun at 1000g in a bench top centrifuge to form 3D spheroids and media topped up to 300 μL per well the next day. Spheroids were grown for 16 days and quantified by cell titre glo viability assays (Promega). To replenish media, 100 μL was removed and replaced with fresh media.

**Mouse xenograft studies**. Nine-week-old female NOD-scid IL2Rγcnull (NSG) and 5-week old male NOD-scid IL2Rγcnull-3/GM/SF (NSGS) mice were obtained from The Jackson Laboratory. Mice were started on –SG or +SG pellets (Test Diet 5CC7) beginning 1 week before cell injections. For HNT34 cells, 104 cells were injected into the flanks of NSG mice on day 0. Mice were continued on +SG or –SG diets, and tumors in the +SG group first became measurable at week 5. Tumor measurements were made weekly for 4 weeks, after which the mice were euthanized. For MUTZ3 cells, 104 cells were injected into the flanks of NSGS mice on day 0, and mice were euthanized at week 4, at which point tumors were dissected from flanks and measured.
Proliferation and 3D growth. Baseline proliferation in MCF-10A and hTERT-IMEC cells was determined in standard DMEM/F12-supplemented media. Media was changed every 2 days, and cells were trypsinized and counted on day 6 (MCF-10A) or day 5 (hTERT-IMECs). For growth factor-independent proliferation, MCF-10A cells were seeded at low density in T25 flasks and grown in standard DMEM/F12-supplemented media, but without EGF for 3 weeks, at which point they were stained with crystal violet and photographed. For mammosphere formation, isogenic MCF-10A cells were grown in standard media combined with Matrigel for 10 days, at which point representative photographs were taken. Soft agar colony formation assays were performed as previously described (56).

Statistics. Significance of ΔPSI of splice junctions between mutant and control MCF-10A cells was tested using a moderated, 2-tailed $t$ test with $P$ less than 0.01 and Benjamini-Hochberg correction $q$ less than 0.05. Genes differentially regulated in SF3B1HET breast tumors from TCGA and METABRIC were included at FDRs of less than 0.10 and less than 0.05, respectively, due to the larger overall number of tumors in the METABRIC cohort. For gene ontology analysis, processes enriched at $P$ less than 0.05 by Fisher’s exact test with $q$ less than 0.05 by Benjamini-Hochberg FDR correction were first captured and then further filtered for those with a fold enrichment of greater than 3. For xenotransplantation, tumor volumes and statistical comparisons were calculated as previously described (58). For growth rescue with PHP, Dunnett’s test was applied to compare control cells with multiple increasing doses of PHP. For all other experiments, a 2-tailed Student’s $t$ test with $P$ less than 0.05 was used to test for significance between groups. All graphs show mean ± SEM. Study approval. The animal care and use protocol of this study was approved by the IACUC of the Johns Hopkins University.

Author contributions

WBD, EH, LPG, DSK, RN, ESC, SD, RDL, A Pandey, J Lauring, and BHP designed research. WBD, EH, NW, LPG, DSK, AR, ESC, BR, LZ, RDL, DS, TG, DJZ, AM, J Lee, AJC, MR, MT, AA, JD, KC, DC, and RMM conducted experiments. WBD, EH, NW, LPG, DSK, AR, RN, ESC, BR, SD, LZ, RDL, DS, TG, AKM, A Pandey, DJZ, AM, J Lee, AJC, JD, AED, KC, DC, RMM, A Patil, PJH, J Lauring, and BHP analyzed data. WBD, ESC, DSK, AKM, A Patil, JD, AED, and BHP wrote the manuscript.

Acknowledgments

This work was supported by grants from the Conquer Cancer Foundation, Breast Cancer Research Foundation, Breast Cancer Now, Ladies Auxiliary of the Veterans of Foreign Wars, Lilly Innovation Fellowship Award, and Department of Defense (W81XWH-17-1-0035) (to WBD); NIH K08HL136894 (to LPG); and the Susan G. Komen Foundation and Abbvie (to BP).

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