




# Defining disease modification in myelofibrosis in the era of targeted therapy

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The development of targeted therapies for the treatment of myelofibrosis highlights a unique issue in a field that has historically relied on symptom relief, rather than survival benefit or modification of disease course, as key response criteria. There is, therefore, a need to understand what constitutes disease modification of myelofibrosis to advance appropriate drug development and therapeutic pathways. Here, the authors discuss recent clinical trial data of agents in development and dissect the potential for novel end points to act as disease modifying parameters. Using the rationale garnered from latest clinical and scientific evidence, the authors propose a definition of disease modification in myelofibrosis. With improved overall survival a critical outcome, alongside the normalization of hematopoiesis and improvement in bone marrow fibrosis, there will be an increasing need for surrogate measures of survival for use in the early stages of trials. As such, the design of future clinical trials will require re-evaluation and updating to incorporate informative parameters and end points with standardized definitions and methodologies. *Cancer* 2022;128:2420-2432. © 2022 The Authors. *Cancer* published by Wiley Periodicals LLC on behalf of American Cancer Society. This is an open access article under the terms of the [Creative Commons Attribution-NonCommercial](#) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes.

**KEYWORDS:** bone marrow fibrosis, disease modification, myelofibrosis, myelofibrosis pathophysiology, targeted therapy.

## A UNIQUE CLINICAL CHALLENGE IN MYELOFIBROSIS

Myelofibrosis (MF) is primarily driven by constitutive activation of the Janus kinase (JAK)/signal transducers and activators of transcription (STAT) pathway. The first approved targeted therapy class for MF, JAK inhibitors (JAKi) have demonstrated amelioration of some key disease symptoms but have otherwise failed to provide survival benefits. This has created a standard for disease response in MF that focuses on symptom relief rather than benefit to progression-free or overall survival (PFS and OS, respectively) or disease modification. This review aims to evaluate, based on clinical and scientific evidence, what constitutes disease modification in MF, together with how this may be measured clinically. The latest developing clinical trial data will be discussed, alongside how these data may inform disease modification, in an attempt to refocus future clinical trial design and MF patient care on potential disease cure and patient survival outcomes.

## INTRODUCTION TO MYELOFIBROSIS

MF pathology includes myeloproliferation, inflammation, bone marrow fibrosis (BMF), extra-medullary hematopoiesis (EMH), splenomegaly, anemia, thrombocytopenia, and constitutional symptoms.<sup>1-3</sup> The median OS of patients with primary MF is ~6 years, and is influenced by a variety of clinical and genetic features including age, mutations, bone marrow fibrosis (BMF), and treatment history.<sup>4-6</sup>

Allogeneic stem cell transplantation (allo-SCT) is currently the only curative option for MF, but its suitability is limited to a minority.<sup>7</sup> The JAKi ruxolitinib (JAK1/2i) and fedratinib (JAK2i) are currently the only approved treatments for patients with MF.<sup>8,9</sup> Despite the efficacy demonstrated by these JAKi in reducing splenomegaly and constitutional

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symptoms across a spectrum of MF patient subgroups,<sup>10-15</sup> JAKi exert little effect on BMF, driver mutation allele frequency (MAF), leukemic transformation or OS. Most patients ultimately experience ruxolitinib failure, leading to treatment discontinuation with limited alternative options.<sup>2,16</sup> There is, therefore, a need for novel treatment options with overt disease-modifying activity.

Greater understanding of MF pathophysiology has unveiled multiple non-JAK targets, leading to the development of several novel agents.<sup>17</sup> However, the lack of an accepted definition and assessment of disease modification is a key hurdle to progressing patient care. In particular, the absence of consistent and standardized parameters by which modification can be assessed limits the impact of emerging data and inadvertently promotes the development of agents that may not provide significant benefit to patients. Furthermore, the lack of coordination across clinical trial end points and clinical practice precludes the inter-trial and -agent comparisons required to optimize treatment pathways.

As with any rapidly evolving field, it is important to acknowledge that many of the clinical trials discussed here are ongoing, and that many of the data that informed our rationale when defining disease modification are immature. The definition we propose will undoubtedly evolve and mature in line with the emergence of future data.

## MYELOFIBROSIS DISEASE COURSE

### *Disease Pathogenesis*

Constitutive activation of the JAK/STAT pathway may be facilitated by driver mutations that confer a fitness advantage.<sup>2</sup> The resultant clonally expanded megakaryocytes cluster in the bone marrow (BM) where they are infiltrated by neutrophils. This results in a “cytokine storm” that generates an inflammatory BM microenvironment, stimulating fibrosis and angiogenesis (Fig. 1; Supporting Table 1).<sup>1,3,6,18,19</sup> As such, BMF is a proximal manifestation of disease biology, stemming from the underlying molecular dysregulation. As a direct result of the dysregulated BM microenvironment, hematopoietic progenitor cells subsequently migrate to sites of EMH, resulting in progressive splenomegaly.<sup>2,20</sup> JAKi alone do not robustly impact BMF, reverse abnormal hematopoiesis, or target the aberrant stem cell niche, suggesting that dysregulated JAK signaling is not the only driving factor of MF disease etiology.

### *Monitoring Disease Progression*

MF has a highly variable disease course, ranging from an indolent, asymptomatic disease, to BM failure or leukemic transformation,<sup>3,17</sup> with somatic mutations playing a central role in determining disease risk.<sup>2,3,21</sup> Progression is

often clinically determined by worsening splenomegaly or leukemic transformation.<sup>13</sup>

Novel, targeted therapies may have the potential to interrupt, or even reverse, the disease trajectory, possibly returning the BM microenvironment to a pre-disease state (Fig. 2).<sup>22,23</sup> However, data are required to inform potential modifiers and to understand the impact of these on PFS and OS. This may serve to accelerate therapeutic developments and facilitate clinical decision-making.

As mentioned, allo-SCT is the only potential cure for patients with MF and thus determination of true remission and positioning of a patient for SCT is an important goal. To this end, measurable residual disease assessment (with the aim of clonal remission) is now recommended in several other hematological malignancies where it forms the backbone of patient management.<sup>24-27</sup> Molecular responses are commonly included in clinical trial designs and may be considered representative of disease modification. As experimental therapies begin to emerge for MF, similar responses may become a new hallmark of treatment and it will be important to define end points and their assessments accordingly. This approach will be of particular significance in the future, when commonly used agents begin to modify disease course and are capable of inducing remission and positioning patients for SCT with greater frequency.

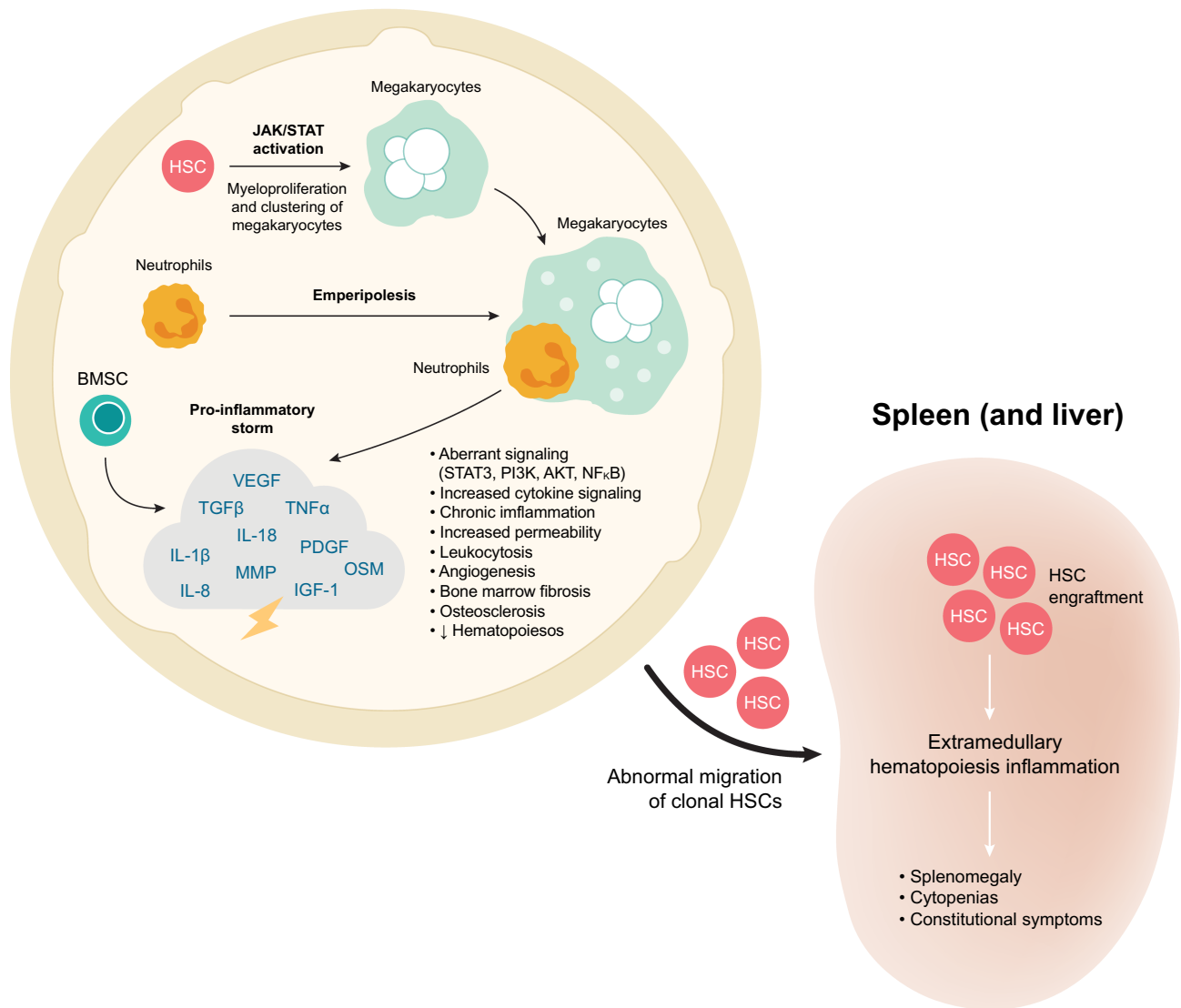
## MYELOFIBROSIS CLINICAL TRIALS

### *Clinical Trial End Points*

Given the role of the JAK/STAT pathway in driving MF pathology, JAK was an obvious initial therapeutic focus. As such, JAKi were the first approved targeted therapy class for MF, and although they have minimal impact on survival, they have been highly effective at controlling splenomegaly and constitutional symptoms. This has resulted in SVR, and total symptom score (TSS) becoming standard end points in MF trials (Supporting Table 2),<sup>11,12</sup> as reflected in the International Working Group for Myelofibrosis Research and Treatment and European Leukemia Net response criteria for myelofibrosis.<sup>13</sup> These symptom and quality-of-life based measures have remained the primary end points for MF treatment trials due to the lack of definitive pathological and/or biochemical criteria to determine MF progression or disease modification. This approach has culminated in a standard for disease response in MF that focuses on symptom relief rather than benefit to PFS, OS, or modification of disease course.

Recently, the number of clinical trial end points has expanded alongside emerging agents, including an increased use of patient reported outcomes (PROs) in parallel with OS, PFS, MAF, cytokine modulation,

## Bone marrow microenvironment



**FIGURE 1.** Overview of disease pathology.<sup>3,6,18,73,77-79</sup> BMSC indicates bone marrow stromal cell; FGF, fibroblast growth factor; HSC, hematopoietic stem cell; IGF-1, insulin-like growth factor-1; IL, interleukin; JAK/STAT, Janus kinase/signal transducer and activator of transcription; MMP, matrix metalloproteinases; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; OSM, oncostatin M; OPG, osteoprotegerin; PDGF, platelet-derived growth factor; PI3K, phosphoinositide 3-kinase; TGF-β, transforming growth factor β; TNF-α, tumor necrosis factor α; VEGF, vascular endothelial growth factor.

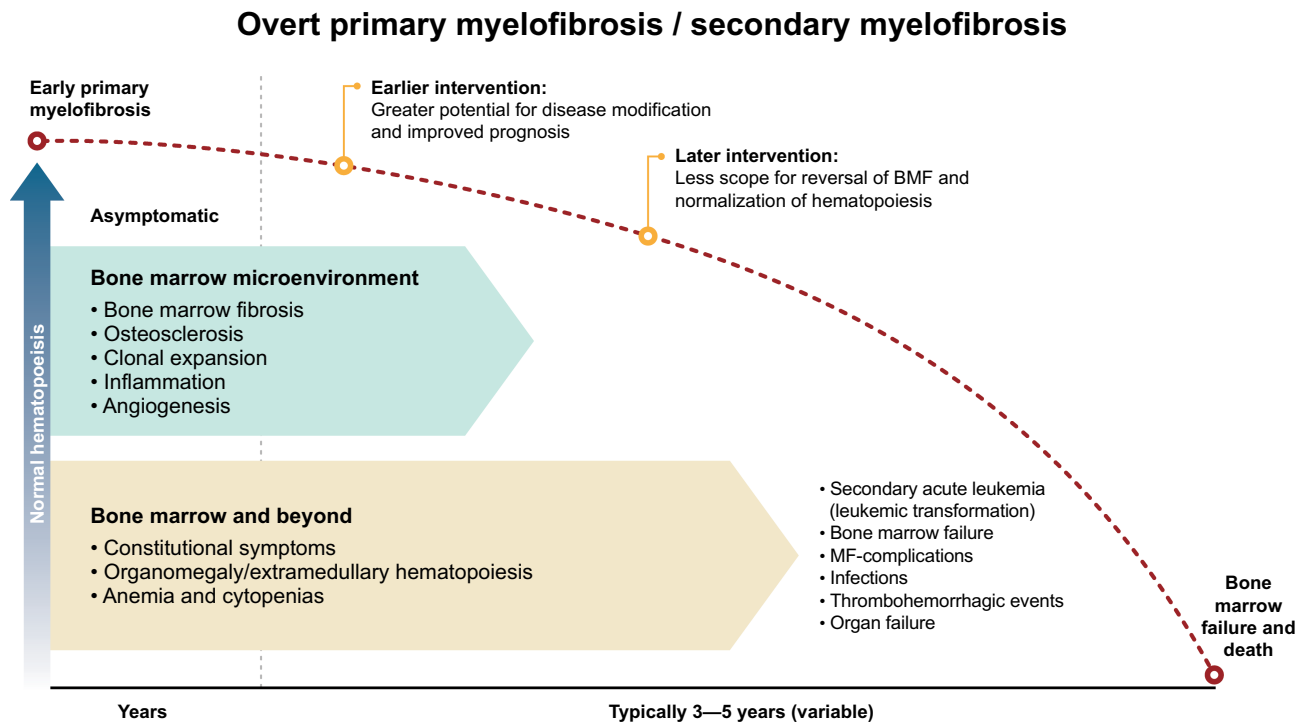
event-free or leukemia-free survival (failure to transform to leukemia), transfusion independence (TI), and reduction in BMF.<sup>28</sup> However, many of these present their own challenges, often due to a lack of standardized definitions.<sup>29-31</sup>

### Latest Clinical Trials of Novel Agents

Here, we discuss the key therapies that begin to inform how disease modification may be defined, with focus on potential disease-modifying effects and the challenges of

defining disease modification. Although many ongoing trials lack novel end points, these should not be overlooked. Importantly, they highlight the need for greater standardization between trial designs as MF research evolves. Early evidence of a progression toward nontraditional primary end points is encouraging,<sup>32,33</sup> and defining key modifiers will help to establish greater uniformity.

An overview of selected targets is presented in **Figure 3**, with the most advanced clinical trial data summarized in **Table 1**. Although a comprehensive analysis



**FIGURE 2.** Natural history of myelofibrosis and potential time points for intervention. The red dotted line represents the decline in normal hematopoiesis along the natural course of disease.<sup>1,2,3,79</sup> BMF indicates bone marrow fibrosis; MF, myelofibrosis.

of all novel therapies is beyond the scope of this review, an overview of the most encouraging clinical trial data to emerge for novel therapies in phase 2/3 development for MF is presented in Supporting Table 3, and Supporting Table 4 lists ongoing trials of promising agents in early development.

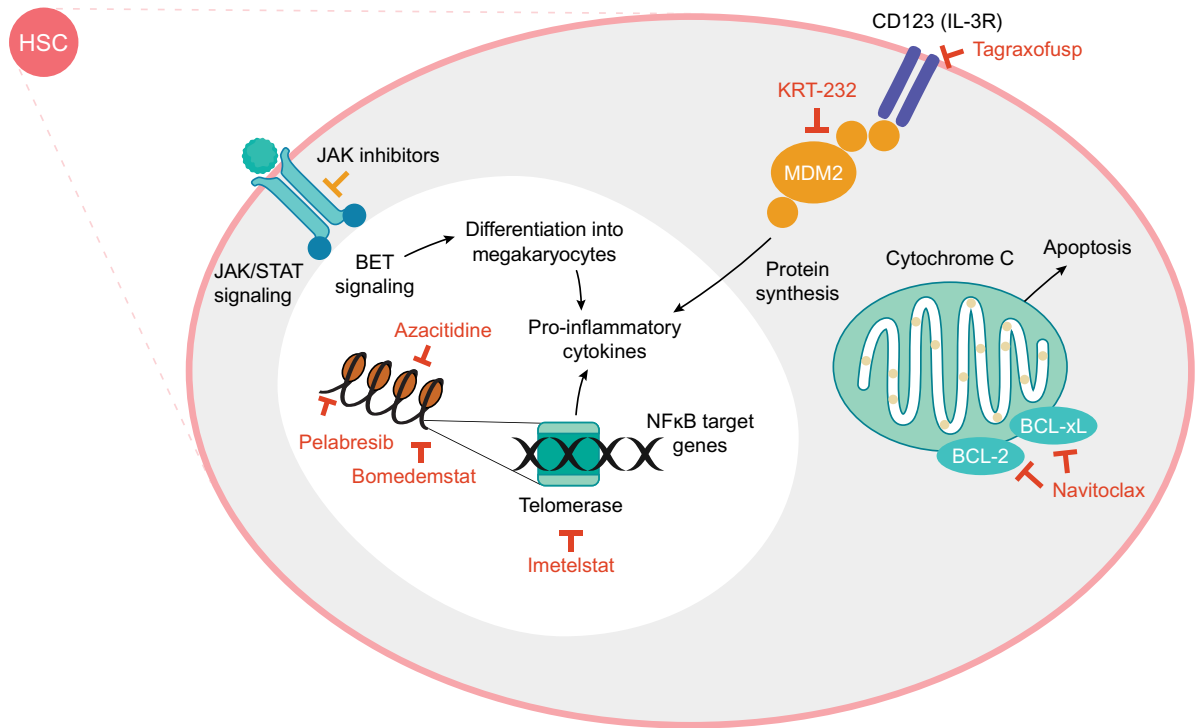
### Pelabresib

MF progenitor cells exhibit altered gene regulation via nuclear factor- $\kappa$ B (NF- $\kappa$ B) pathway activation that may sustain the inflammation associated with disease progression and transformation via aberrant cytokine signaling.<sup>34,35</sup> Bromodomain and extra-terminal motif inhibitors (BETi) have been developed to exploit this, with the aim of attenuating NF- $\kappa$ B signaling and suppressing cytokine release.<sup>36</sup> Pelabresib is the most advanced BETi, with recruitment currently underway in phase 2 and 3.<sup>37-39</sup> The phase 2 trial, MANIFEST, included 3 arms: 1) pelabresib as monotherapy in JAKi-experienced patients; 2) as “add-on” to ruxolitinib in patients with inadequate response to ruxolitinib; and 3) in combination with ruxolitinib in JAKi-naive patients. Interim data from MANIFEST have demonstrated improvement in BMF of  $\geq 1$  grade in 33% of patients across the arms (21%, 41%, and 33%

in arms 1, 2, and 3, respectively).<sup>40</sup> Translational studies have also reported broad clinical responses regardless of baseline mutational status, with pelabresib treatment associated with significant reduction of several cytokines in ruxolitinib-naive and ruxolitinib-experienced patients, increased erythroid progenitors, and improved megakaryocyte histology. With survival data yet to be reported, these data suggest disease-modifying potential.<sup>40,41</sup> The primary trial end point,  $\geq 35\%$  SVR from baseline (SVR35) at week 24, was achieved by 24%, 21%, and 67% of patients, respectively.<sup>42-44</sup> This suggests that potentially disease-modifying activity may be largely independent of prior therapy or treatment order, unlike SVR that demonstrated an association with treatment history. Although any definitive association between reduced BMF and disease modification remains to be characterized, these data are supportive of BMF as a potential indicator of disease modification.

The encouraging activity reported from JAKi-naive patients in MANIFEST support the assessment of JAKi combination therapy in the first-line setting and will be further explored in MANIFEST-2, where JAKi-naive patients will be randomized to pelabresib plus ruxolitinib or placebo plus ruxolitinib. Bone marrow

## Hematopoietic stem cell (HSC)



- ⊥ Agents with potential disease-modifying activity
- ⊥ Currently approved agents

**FIGURE 3.** Novel and potentially disease-modifying therapeutic targets in myelofibrosis.<sup>80-82</sup> BCL indicates B-cell lymphoma; BET, bromodomain and extra-terminal motif; IL, interleukin; JAK/STAT, Janus kinase/signal transducers and activators of transcription; MDM2, mouse double minute 2; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; R, receptor.

morphology and proinflammatory cytokine modulation will also be explored alongside the primary end point of SVR35.<sup>38</sup>

### Bomedemstat

Hematopoiesis is dependent on the epigenetic modifier lysine-specific demethylase 1 (LSD1), which has a specific role in megakaryocyte maturation.<sup>45</sup> Bomedemstat was developed as an irreversible inhibitor of LSD1 and is under investigation in the phase 1/2 setting. Interim data have reported improvement in BMF of  $\geq 1$  grade for 17% of evaluable patients and reductions in MAF in driver and HMR mutations in 42% of patients. MAF reduction was found to correlate with improvements in spleen volume and/or TSS.<sup>46</sup> Although any correlation with survival end points remains to be determined, these data further support the need for BMF and/or mutation-focused short-term end points to inform their utility in defining disease modification. Serving to

bolster this assertion, spleen volume and TSS reduction or stability were also observed in the vast majority of these patients.

### Navitoclax

Targeting the B-cell lymphoma-2 (BCL-2) antiapoptotic pathway has been remarkably successful in chronic lymphocytic leukemia and acute myeloid leukemia.<sup>47,48</sup> Preclinically, *JAK2V617F* mutated CD34+ HSC exhibit apoptotic resistance through overexpression of BCL-2 family proteins.<sup>49,50</sup> Navitoclax is a BCL-2/BCL-xL inhibitor that is undergoing phase 2 investigation as monotherapy and in combination with ruxolitinib in the REFINE study. Interim analyses have demonstrated improvement in BMF of  $\geq 1$  grade for 33% (11/33) of patients and  $>10\%$  reduction in driver gene MAF in 46% (12/26) of patients. The clinical impact of these data remains to be determined—median OS was not reached at a median follow-up

**TABLE 1.** Novel Agents in Development for MF With Potential for Disease-Modifying Activity

Study Name/No.	Drug	MoA	Phase/Status	Population	Comparator	Primary End Point	BMF	Mutation Burden	Survival	Notes
Epigenetic modulation NCT02158858 (MANIFEST) <sup>40,41,43,44,63,74</sup>	Pelabresib ± Rux	BET ± JAK1/2 inhibition	1/2 Ongoing	Arm 1: JAKi-experienced (pelabresib) Arm 2: JAKi-experienced (pelabresib + Rux) Arm 3: JAKi-naive (pelabresib + Rux) N ≈ 271	Placebo + Rux None	Arm 1: SVR35 at wk 24: 24% TD → TI: 21% Arm 2: SVR35 at wk 24: 21% TD → TI: 36% Arm 3: SVR35 at wk 24: 67%	1 Grade improvement: 33%	NR	NR	Several cytokines suppressed with mono and combination therapy
NCT04603495 (MANIFEST-2) <sup>38</sup> NCT03136185 <sup>46</sup>	Pelabresib + Rux Bomedemstat	BET ± JAK1/2 inhibition LSD1 inhibitor	3 Ongoing 2 Ongoing	JAKi- and BETi-naive, N ≈ 310 Rux-naive and -experienced, N = 89	None	SVR35 at wk 24 AEs: 89%	1 Grade improvement: 17% (stable 66%)	Reduction in MAF in driver and HMR mutations: 42% (stable in 44%)	NR	MAF reduction correlated with SVR and/or TSS
NCT01787487 <sup>61</sup>	Azacitidine + Rux	HMA + JAK1/2 inhibition	2	Rux-naive, azacitidine-naive, N = 60	None	ORR: 74%	Improvement in BM morphology: 61%	NR	Median OS: not reached	
Hematopoietic stem cell compartment/bone marrow microenvironment NCT03662126 (BOREAS) <sup>60</sup>	Navtemadlin	MDM2 inhibition	2/3 Ongoing	JAKi-experienced, N = 113	None	NR	≥1 Grade improvement: 27%	Reduction in MAF ≥20%: 34% (complete reduction 29%)	NR	Reduced circulating CD34+ cells and TNF-α; MAF, BMF, CD34+, and TNF-α correlated with SVR and/or TSS
NCT04640532	Navtemadlin ± TL-895	MDM2 inhibition + tyrosine kinase inhibition	1/2 Ongoing	JAKi-experienced, N ≈ 116	None	MTD/MAD; RP2D; SVR35 at wk 24	—	—	—	
NCT04485260 <sup>69</sup>	Navtemadlin + Rux	MDM2 inhibition + JAK1/2 inhibition	1b/2 Ongoing	Rux-experienced, N ≈ 78	None	RP2D	—	—	Secondary end points: OS, PFS, LFS	
NCT02268253 <sup>63</sup>	Tagraxofusp	Anti-CD123	1/2 Ongoing	JAKi-experienced, N ≈ 130	None	AEs; RR: 60% SD	NR	NR	Median OS: 31 mo	
Apoptosis NCT03222609 (REFINE) <sup>31,51</sup>	Navitoclax ± Rux	BCL inhibition ± JAK1/2 inhibition	2 Ongoing	Rux-experienced, N ≈ 174	None	SVR35 at wk 24: 27% (dual therapy)	≥1 Grade improvement: 33%	>10% reduction in driver gene MAF: 46%	Median OS: not reached	Changes in MF-associated cytokines were correlated with SV changes
NCT04472598 (TRANSFORM-1) <sup>52</sup>	Navitoclax + Rux	BCL inhibition + JAK1/2 inhibition	3 Ongoing	JAKi-naive, N ≈ 230	Placebo	SVR35 at wk 24	Secondary end point	—	Additional end points: OS, LFS, PFS	

TABLE 1. Continued

Study Name/No.	Drug	MoA	Phase/Status	Population	Comparator	Primary End Point	BMF	Mutation Burden	Survival	Notes
NCT04468984 (TRANSFORM-2) <sup>63</sup>	Navitoclax + Rux	BCL inhibition + JAK1/2 inhibition	3 Ongoing	JAKi-experienced, N ≈ 330	BAT	SVR35 at wk 24	Secondary end point	—	Additional end points: OS, LFS, PFS	
Telomerase NCT04576156 (MYF3001)	Imetelstat	Telomerase inhibition	3 Ongoing	JAKi-experienced, N ≈ 320	BAT	OS	Secondary end point	—	—	Biomarker and mutation analyses will be performed
NCT02426086 (IMbark) <sup>65</sup>	Imetelstat	Telomerase inhibition	2 Complete	JAKi-experienced, N = 59	None	SVR35 at wk 24: 10.2% ≥50% reduction in TSS at wk 24: 32.2%	Improvement: 40.5%	Reduction in driver mutation MAF: 42.1%	Median OS: 29.9 mo	
Immune therapies NCT01178281 (RESUME) <sup>75</sup>	Pomalidomide	IMiD	3 Complete	N = 252	Placebo	RBC-TI ≥84 days within 6 mo: 16% vs 16% (P = 1.00)	NR	NR	NR	
NCT01644110 (POMINC/MPNSG-0212) <sup>76</sup>	Pomalidomide + Rux	IMiD + JAK1/2 inhibition	1/2 Ongoing	Rux-naive and -experienced, N ≈ 90	None	CI, 18% (low dose pom); 20% (high dose pom)	NR	NR	NR	
NCT03069326 <sup>87</sup>	Thalidomide + Rux	IMiD + JAK1/2 inhibition	2 Ongoing	Rux-experienced, N ≈ 65	None	ORR, 60%	NR	NR	NR	Significant increase in platelet count after cycle 3 vs baseline (P < .01); platelet count increase in 75% of patients with thrombocytopenia

Abbreviations: ACVR, activin A receptor; AE, adverse event; BAT, best available therapy; BCL, B-cell lymphoma; BET, bromodomain and extra-terminal; BM, bone marrow; BMF, bone marrow fibrosis; CI, clinical improvement; CRP, C-reactive protein; ELN-IWG, European LeukemiaNet international working group; Hb, hemoglobin; HMA, hypomethylating agent; HMR, high molecular risk; HR, hazard ratio; IL, interleukin; IMiD, immunomodulatory; IRAK, interleukin 1 receptor associated kinase; JAK, Janus kinase; JAKi, JAK inhibitor; LFS, leukemia-free survival; LSD, lysine-specific histone demethylase; MAD, maximum administered dose; MAF, mutant allele frequency; MDM2, mouse double minute 2 homolog; MF, myelofibrosis; MoA, mechanism of action; MTD, maximum tolerated dose; ORR, overall response rate; OS, overall survival; PD-1; programmed cell death protein 1; PFS, progression-free survival; PI3K, phosphoinositide 3-kinase; NR, not reported; QD, once daily; QW, once weekly; RBC, red blood cell; RP2D, recommended phase 2 dose; Rux, ruxolitinib; SD, stable disease; SVR35, ≥35% spleen volume reduction from baseline; TD, transfusion dependence; TGF, transforming growth factor; TNF-α, tumor necrosis factor alpha; TI, transfusion independence; TSS, total symptom response

of 105 weeks—and again reflects the need for further understanding and standardization of such end points. Alongside these, SVR35 at any time was achieved by 44% of patients (27% at week 24), which was durable regardless of HMR.<sup>31,51</sup> Furthermore, direct correlations were observed between several MF-associated cytokines and changes in spleen volume.<sup>31</sup> Phase 3 recruitment is now ongoing to investigate navitoclax plus ruxolitinib versus placebo plus ruxolitinib in JAK2i-naïve patients (TRANSFORM-1)<sup>52</sup> and versus best available therapy (BAT) in JAK2i-experienced patients (TRANSFORM-2). The traditional primary end point of SVR35 will be explored alongside additional analyses of BMF and time-to-event survival measures.<sup>53</sup>

### Imetelstat

Telomerase is upregulated in many cancers, and CD34+ hematopoietic cells in MPNs are characterized by shortened telomeres.<sup>54</sup> Preclinical models of the competitive telomerase inhibitor, imetelstat, have demonstrated selective inhibition of pre-leukemic stem cell transformation via downregulation of hTERT and decreased ADAR1 activity.<sup>55</sup> In the phase 2 study, IMbark, 41% of patients experienced reversal of BMF and 42% had reduced MAF of driver mutations, both of which correlated with improved OS. Median OS was 29.9 months.<sup>56</sup> Cytogenetic analyses within IMbark have demonstrated the selective targeting of malignant cells by imetelstat, further supporting the potential for disease-modifying activity.<sup>57</sup> Interestingly, improvements in the primary end points of SVR35 and TSS at week 24 were modest (10% and 32%, respectively),<sup>56</sup> and not correlated BMF or MAF, clearly presenting the need for more informative end points to be adopted across MF clinical trials. Imetelstat is now under further evaluation versus BAT for patients refractory to JAKi in the phase 3 trial MYF3001, which will evaluate impact on malignant clones leading to disease modification alongside a primary end point of OS. This deviation from traditional end points reflects the reform required across clinical trials to better understand the full potential of emerging therapies.

### Navtemadlin

Inhibition of mouse double minute 2 (MDM2) drives selective depletion of *JAK2V617F* mutated stem cells.<sup>37,58</sup> Navtemadlin, a potent MDM2 inhibitor is undergoing assessment as monotherapy and combination therapy.<sup>59</sup> Interim data from the phase 2/3 BOREAS trial of navtemadlin monotherapy in JAKi relapsed/refractory

(R/R) patients are encouraging, with improved BMF observed in 27% (12/45) of patients and stable scores in 51%. Best driver gene reduction  $\geq 20\%$  was reported in 34% (22/65) of patients and 29% (19/65) had a complete MAF reduction below the limit of detection. Furthermore, navtemadlin was associated with reduced levels of circulating CD34+ cells and tumor necrosis factor  $\alpha$  (TNF- $\alpha$ ). Each of these parameters correlated with SVR, and an association was also observed between fibrosis scores and mutational burden.<sup>60</sup> As navtemadlin is investigated in phase 3, correlations between these potentially disease-modifying parameters and survival outcomes are eagerly awaited.

### Azacitidine

Azacitidine is a hypomethylating agent currently undergoing testing as “add-on” to ruxolitinib in JAKi-naïve patients. Although interim data are encouraging in this single arm study, with improvement in BM morphology reported for 61% of patients, and an overall response rate of 74%, the additional benefit of azacitidine remains to be fully determined. However, it is noteworthy that median OS had not been reached after a median follow-up of 35 months.<sup>61</sup>

### Tagraxofusp

Tagraxofusp is CD123-directed cytotoxin selected for testing in MF as CD123 is an established marker of leukemic stem cells.<sup>62</sup> Interim results of a phase 2 study were modest, with spleen responses observed in 45% of patients at 24 weeks and a median OS of 31 months, the latter of which may prove indicative of disease-modifying activity.<sup>63</sup>

### Immune therapies

Allo-SCT is an immune therapy that represents the only definitively disease-modifying option currently available, and successful transplant generally leads to the reversal of BMF.<sup>64</sup> The atypical and dysregulated immune environment associated with MF provides a variety of additional immune targets. Interferon (IFN)-based regimens have shown promise, with a survival benefit of pegylated IFN- $\alpha 2$  demonstrated in the long-term follow-up of patients with intermediate or high-risk disease, whereby the median OS of 89 months was longer than expected and accompanied by a reduction in *JAK2V617F* burden.<sup>65,66</sup> Additional immune therapies including immunomodulatory drugs and checkpoint inhibitors are undergoing investigation in combination with ruxolitinib.<sup>67,68</sup>



Finally, the JAK/STAT pathway remains an attractive target and several JAKi are in development. Although there is no evidence to suggest these modify disease course, JAKi are ideal candidates for combination therapy and many trials are examining novel agents in combination with ruxolitinib.

Readers should exercise caution when interpreting the above data, as many of these trials are ongoing; data are preliminary and often based on small numbers with incomplete follow-up. However, these early reports are encouraging, provide initial rationale for defining disease modifying parameters, and hold promise that significant evolution of the MF treatment landscape is on the horizon.

### PROPOSED DEFINITION OF DISEASE MODIFICATION IN MYELOFIBROSIS

The current benchmark for curative therapy in MF is allo-SCT, which replaces disease through healthy repopulation of the stem cell compartment. When considering disease modification, a treatment is not expected to replace disease, but rather alter the disease biology, with the aim of reversing disease trajectory. As such, true disease modification is difficult to measure without standardized assessment or consensus recommendation to guide clinical evaluation. This makes it critical for any definition of disease modification to represent true modifiers and mechanisms of improvement, rather than resultant downstream effects.

With consideration of the current knowledge and the rationale provided by available clinical trial data, we propose the following definition of disease-modifying activity (Table 2):

Disease modification in MF is defined as therapy that exerts a clinically meaningful impact on survival outcomes and/or restoration of normal hematopoiesis in conjunction with improvement in bone marrow fibrosis through a substantial and durable reduction in the clonal burden of disease.

It seems inevitable that achieving disease modification will also lead to beneficial downstream effects such as the elimination of symptoms and splenomegaly and improved PROs.

### DISCUSSION AND FUTURE PERSPECTIVES

The treatment landscape of MF has remained almost static for a decade but is set to evolve rapidly as understanding of the molecular pathogenesis of MF sheds light on novel

therapeutic targets and the possibility of selectively depleting the malignant HSC compartment (Fig. 3).<sup>17</sup>

As novel treatment strategies emerge, their optimal use and place in the treatment paradigm will need to be determined. Ascertaining the appropriate timing of interventions to maximize the potential for disease modification will be key. Trials tend to take place in the heavily pretreated setting; however, it is logical that the greatest impact of disease modifying treatment will be observed when initiated early, before clonal evolution (Fig. 2).

Despite the limited disease-modifying activity of JAKi, the JAK/STAT pathway remains a pivotal feature of MF pathology and it is unlikely that JAK inhibition will be relinquished. Rather, synergy between inhibitors of JAK and non-JAK targets may positively impact disease modification, optimizing clinical responses. Treatment strategies that combine JAKi with novel agents, especially given the current reliance on JAKi to control disease symptoms, are likely to feature heavily as clinical trial programs develop.

Several agents and combinations undergoing study in MF are suggestive of disease modification. Although interpretation across trials is limited by the lack of control arms and disparities in the definition and recruitment of patients with ruxolitinib failure, together with inconsistent trial designs, available data are beginning to inform recommendations for defining and measuring disease-modifying activity. Emerging data from novel end points will facilitate this evolution and we await the correlative assessments between modifiers and OS. Future trials that can demonstrate significant correlation between reduction in BMF grade or reduction in clonal disease burden with increased median PFS or OS will provide evidence for the first wave of MF treatments that modify disease.

These exciting developments highlight the need for new surrogate measures and study end points, particularly for use in the early stages of trials before survival read outs, to help identify the most promising new approaches and accelerate their approval. As the landscape evolves, it will be important to standardize such parameters across clinical trials to better define the potential for disease modification and facilitate inter-trial comparability. Here, we present our recommendation for how disease modification of MF should be defined, informed by current knowledge and available data.

Looking to the future, assessment of disease burden within the HSC compartment may emerge, not only as a defining end point of disease modification, but a measure

**TABLE 2.** Proposed Parameters, With Rationale, Considered as Evidence of Disease Modification

Parameters	Rationale	Supporting Data From Novel Agents	Limitations
<p>Primary outcomes of disease modification</p> <p>OS</p> <p>Event-, progression- or leukemia-free survival</p>	<ul style="list-style-type: none"> <li>Most critical outcomes of any life-threatening disease treatment</li> </ul>	<ul style="list-style-type: none"> <li>Not generally a primary end point, survival outcomes have mostly yet to be reported</li> <li>Imetelstat (IMbark) and tagraxofuspup (NCT02268253) reported median OS of 30 and 31 mo, respectively<sup>56,63</sup></li> </ul>	<ul style="list-style-type: none"> <li>Require lengthy follow-up that may not be compatible with timelines for drug approval</li> </ul>
<p>Key modifiers</p> <p>BMF</p>	<ul style="list-style-type: none"> <li>BMF grade directly influences OS, is the most proximal result of the clonal malignancy and directly contributes to extra-medullary hematopoiesis, splenomegaly, insufficient blood cell production, anemia, and thrombocytopenia<sup>69,70</sup></li> <li>Successful allo-SCT generally leads to the reversal of BMF<sup>64</sup></li> </ul>	<p>Improved BMF and correlation to OS reported by:</p> <ul style="list-style-type: none"> <li>Imetelstat (IMbark)<sup>56</sup></li> <li>BMF improvement demonstrated by:                             <ul style="list-style-type: none"> <li>Pelabresib (MANIFEST)<sup>40</sup></li> <li>Bomedemstat (NCT03136185)<sup>46</sup></li> <li>Navitoclax (REFINE)<sup>31</sup></li> <li>Navtemadlin (BOREAS)<sup>60</sup></li> </ul> </li> <li>Correlation with survival outcomes remain to be determined</li> </ul>	<ul style="list-style-type: none"> <li>No clear definition or guidelines regarding the time points of sequential measurements and grading of fibrosis</li> <li>Significance of reduced BMF is unclear unless associated with improvement of cytopenias</li> </ul>
<p>Clonal disease/mutational burden</p>	<ul style="list-style-type: none"> <li>Clonal disease burden is correlated with disease phenotype and evolution is thought to be predictive of patient outcomes<sup>71-73</sup></li> </ul>	<p>Reduced MAF and correlation to OS reported by:</p> <ul style="list-style-type: none"> <li>Imetelstat (IMbark)<sup>56</sup></li> </ul> <p>Reductions in MAF reported by:</p> <ul style="list-style-type: none"> <li>Bomedemstat (NCT03136185)<sup>46</sup></li> <li>Navitoclax (REFINE)<sup>31</sup></li> <li>Navtemadlin (BOREAS)<sup>60</sup></li> </ul> <p>Correlation with survival outcomes remain to be determined</p>	<ul style="list-style-type: none"> <li>No data or guidelines regarding what might be considered a clinically meaningful reduction in mutational burden and association with survival outcomes</li> <li>Need to define how this is measured with regard to driver mutations, additional mutations and clonal hematopoiesis</li> </ul>
<p>Cytokine modulation</p>	<ul style="list-style-type: none"> <li>The MF inflammatory cytokine signature is a consequence of the malignant clone and a key modifier of the bone marrow microenvironment and promoter of malignant hematopoiesis<sup>18,19</sup></li> </ul>	<p>Reductions in key cytokine expression demonstrated by:</p> <ul style="list-style-type: none"> <li>Pelabresib (MANIFEST)<sup>40</sup></li> <li>Navitoclax (REFINE)<sup>31</sup></li> <li>Navtemadlin (BOREAS)<sup>60</sup></li> </ul>	<ul style="list-style-type: none"> <li>Inadequate data regarding the association of cytokine modulation with survival outcomes</li> </ul>

Abbreviations: allo-SCT, allogeneic stem cell transplant; BMF, bone marrow fibrosis; MAF, mutation allele frequency; OS, overall survival.

that begins to align MF with other hematological malignancies in which disease modification is already established. Additionally, baseline characteristics or genetic factors that may impact the potential for disease modification will need to be identified and understood in terms of what may be achieved.

In summary, the possibility of disease modification has the potential to revolutionize clinical practice and treatment decision-making for patients with MF. As novel end points begin to emerge, it will be important to re-evaluate clinical trial designs, and potentially redefine disease modification, adding new end points to survival outcomes, to ensure the true potential for disease modification and MF therapy is realized. Standardized definitions and assessments are needed across clinical trials, along with the inclusion of patients with newly diagnosed disease, where the greater potential for disease modification may lie.

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#### CONFLICT OF INTEREST DISCLOSURE

Naveen Pemmaraju had consulting or advisory roles with Celgene, Stemline, Incyte, Novartis, Mustang Bio, Roche Diagnostics, and LFB; received honoraria from Celgene, Stemline, Incyte, Novartis, Mustang Bio, Roche Diagnostics, and LFB; received grants and/or funding from Affymetrix and Sager Strong Foundation; and had board memberships (noncompensated) with Dan's House of Hope (board of directors) and HemOnc Times/Oncology Times (board member and editor-in-chief). Srdan Verstovsek had consulting or advisory roles with Celgene, Constellation Pharmaceuticals, Incyte, Novartis, Pragmatist, and Sierra and received research funding from Blueprint Medicines, Celgene, CTI BioPharma Corp, Genentech, Gilead Sciences, Incyte, Novartis, NS Pharma, Promedior, and Roche. Ruben Mesa had consulting roles with Constellation, La Jolla Pharmaceutical Company, Novartis, Pharma, and Sierra Oncology and received research funding (institutional) from AbbVie, Celgene, Constellation, CTI, Genotech Pharma, Incyte, Promedior, and Samus. Vikas Gupta provided consultancy for BMS-Celgene, AbbVie, Pfizer, Roche, Constellation Pharma, Sierra Oncology, and Novartis; served on advisory committees for BMS-Celgene, Sierra Oncology, and Novartis; and received research funding from Incyte and Novartis. Jacqueline S. Garcia had consulting/advisory roles with AbbVie, Takeda, and Astellas and received research funding (institutional) from AbbVie, Genentech, Pfizer, Prelude, and Astra Zeneca. Joseph M. Scandura served on consulting/advisory boards for AbbVie, Constellation, and SDP Oncology and received research support from AbbVie and Constellation. Stephen T. Oh served on consulting/advisory boards for Novartis, Kartos Therapeutics, CTI BioPharma, Celgene/Bristol-Myers Squibb, Disc Medicine, Blueprint Medicines, PharmaEssentia, Constellation, Geron, AbbVie, Sierra Oncology, and Incyte. Francesco Passamonti had advisory board roles with AbbVie, BMS, Novartis, Janssen, and Astellas and received speaker fees from AbbVie, BMS, and Novartis. Konstanze Döhner reported consulting/advisory roles with and honoraria from AbbVie, Celgene/BMS, Novartis, CTI BioPharma Corp, and Roche. Adam J. Mead received honoraria for consulting and speaker fees from Novartis, Celgene/BMS, AbbVie, CTI, Sierra Oncology, Karyopharm, Sensyn, Incyte, Galecto, Pfizer and Gilead; received research funding from Celgene/BMS, Novartis, and Galecto; and is a cofounder and equity holder in Alethiomics, Ltd, a spin-out company from the University of Oxford.

#### AUTHOR CONTRIBUTIONS

All authors had access to relevant data and participated in the writing, review, and approval of the manuscript.

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