Novel therapeutics in myeloproliferative neoplasms

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Abstract
Hyperactive signaling of the Janus-Associated Kinase/Signal Transducers and Activators of Transcription (JAK/STAT) pathway is central to the pathogenesis of Philadelphia-chromosome-negative myeloproliferative neoplasms (MPN), i.e., polycythemia vera (PV), essential thrombocythemia (ET), and primary myelofibrosis (PMF) which are characterized by inherent biological and clinical heterogeneity. Patients with MPNs suffer from substantial symptom burden and curtailed longevity due to thrombohemorrhagic complications or progression to myelofibrosis or acute myeloid leukemia. Therefore, the management strategies focus on thrombosis risk mitigation in PV/ET, alleviation of symptom burden and improvement in cytopenias and red blood cell transfusion requirements, and disease course alteration in PMF. The United States Food and Drug Administration’s (USFDA) approval of two JAK inhibitors (ruxolitinib, fedratinib) has transformed the therapeutic landscape of MPNs in assuaging the need for frequent therapeutic phlebotomy (PV) and reduction in spleen and symptom burden (PV and PMF). Despite improving biological understanding of these complex clonal hematopoietic stem/progenitor cell neoplasms, none of the currently available therapies appear to modify the proclivity of the disease per se, thereby remaining an urgent unmet clinical need and an ongoing area of intense clinical investigation. This review will highlight the evolving targeted therapeutic agents that are in early- and late-stage MPN clinical development.

Keywords: MF, ET, PV, JAK-STAT, CALR, Ruxolitinib, Fedratinib, Pacritinib, Imetelstat, CALR vaccine

Introduction
Polycythemia vera (PV), essential thrombocythemia (ET), and primary myelofibrosis (PMF) are clonal myeloproliferative neoplasms (MPN) with distinct hematological and clinicopathologic features that can be viewed as a disease spectrum [1]. Approximately 90% of patients with MPNs harbor mutations involving the JAK2, CALR, or MPL genes (phenotypic drivers in MPN), resulting in hyperactivation of the Janus-Associated Kinase/Signal Transducers and Activators of Transcription (JAK/STAT) signaling pathway [2–4]. Additionally, they may harbor mutations in the epigenetic modifiers (DNMT3A, TET2, ASXL1, IDH1/2, EZH2), RNA splicing (SRSF2, U2AF1), tumor suppressor (TP53) genes that co-operate with each other, and the driver mutations to bestow a more advanced disease phenotype [5]. While patients with PMF suffer from debilitating constitutional symptoms, progressive splenomegaly and cytopenias, PV and ET patients experience microvascular symptoms (headaches, erythromelalgia, Raynaud syndrome) and grievous life-threatening thromboses (arterial and venous) [6, 7]. In general, patients with MPN are at an increased risk of developing thrombosis compared to the general population (PV > ET > PMF) [8] and may progress to acute myeloid leukemia (PMF > PV > ET) [9]. Therefore, prevention of thrombosis and disease progression form the two-pronged approach in the treatment strategy of MPNs. While the current prognostic models in PV and ET are predicated on the clinical and hematological parameters...
that predict the risk of recurrent thrombosis [10, 11], the integration of molecular and clinical data in PMF has allowed for more refined risk stratification and early evaluation for hematopoietic cell transplantation (HCT) [12], which remains the only curative treatment modality.

**Agents in clinical development in PV and ET**

Age (<60 vs. >60 years) and the history of thrombosis form the basis of the risk-adapted approach informing the management decisions in PV and ET as thrombosis is the leading cause of preventable death in these MPNs [11, 13]. Patients younger than 60 years of age with no history of thrombosis are categorized as “low risk” and managed conservatively with therapeutic phlebotomy to maintain a hematocrit less than 45% in PV [14]. In both PV and ET, these low-risk patients are counseled to optimize cardiovascular risk factors (smoking, blood pressure, obesity) and prescribed low-dose aspirin for thrombosis prevention except in JAK2 wild-type ET patients who are deemed as very low risk and maintained on observation only [15]. Cytoreductive therapy is reserved for those patients with “high-risk” features in both ET and PV and low-risk PV patients suffering from uncontrolled symptoms, symptomatic splenomegaly, and intolerance to therapeutic phlebotomy. While hydroxyurea (HU) is the initial drug of choice [16], pegylated interferon-α (IFNα) is preferred in younger patients desiring offspring as HU is a potential teratogen [18, 19] and ruxolitinib, a JAK1/2 inhibitor for those PV patients who are intolerant or resistant to HU [20, 21].

Although low-risk PV patients are managed with therapeutic phlebotomy and aspirin, they still experience higher than normal rates of thrombosis compared to the general population [8] as they may not have a well-controlled hematocrit (<45%) between visits that may predispose them to poor outcomes secondary to hyperviscosity. Additionally, therapeutic phlebotomy leads to iron deficiency-related symptoms that may exacerbate/mimic PV-related symptoms. In this regard, hepcidin mimetics are being evaluated in PV as an alternative to therapeutic phlebotomy as hepcidin regulates iron metabolism, limits intestinal iron absorption, and restricts erythroid differentiation. Preclinical studies of minihepcidin in murine models of PV have shown that prolonged administration curbs the availability of iron to erythroid precursors, thereby impeding erythropoiesis, resulting in the normalization of hematocrit [22]. PTG300, a self-injectable hepcidin mimetic administered weekly, is currently being evaluated as a “medical phlebotomy agent” in a phase II study in PV patients requiring frequent therapeutic phlebotomy (NCT04057040).

Given the inherent risk of thrombosis in PV regardless of the current risk stratification, the move to initiate cytoreductive therapy to mitigate the risk of thrombosis in low-risk PV patients is gaining momentum. IFNα may have disease-modifying activity in MPNs as evidenced by preclinical studies; several small phase 2 studies have shown that IFNα treatment can induce molecular and cytogenetic responses in treated MPN patients, although the results vary according to the series. While some investigators have reported that patients harboring TET2 co-mutations do not respond as well to IFNα as those harboring wildtype, others have reported that patients with low JAK2 V617F variant allele frequency (VAF) are more likely to achieve complete hematological response with IFNα treatment than those with higher VAF at baseline [23–26]. Nevertheless, IFNα appears to induce hematological, molecular, and cytogenetic responses [27] and the clinical benefit of IFNα appears to be optimal when employed earlier in the disease course. In this context, the LOW PV trial is evaluating Ropeginterferon alfa-2b (Ropeg) compared to therapeutic phlebotomy in a phase II randomized clinical trial in low-risk PV patients. Ropeg is a monopegylated interferon that overcomes the shortcomings of IFNα (administered weekly), allowing for less frequent dosing (administered every two weeks) and improved patient tolerability ensuring long-term patient compliance [28]. Maintaining hematocrit ≤45% for 12 months in the absence of progressive disease is the primary composite endpoint of the LOW PV trial. The recently presented preplanned interim analysis shows that 84% of patients on the ropeg arm achieved the primary composite endpoint (60% in the phlebotomy arm; OR=3.5, 95% CI 1.3–10.4, p=0.008) with a lower number of required therapeutic phlebotomy after one year of treatment (43% vs. 57%; p=0.024). Ropeg was well tolerated with no significant difference in adverse events (AE) between both treatment arms. This trial has stopped enrollment in view of the resounding efficacy, and follow-up will continue for 2 years per protocol [29]. The final results of this trial may have practice-changing implications in patients with low-risk PV.

Previously, the Myeloproliferative Disorders Research Consortium (MPD-Rc) 112 study and MPD-Rc 111 study have highlighted the activity of IFNα in treatment-naïve and HU-resistant/refractory ET/PV patients, respectively [26, 30]. Most recently, phase III randomized controlled trials, PROUD-PV and its extension study CONTINUATION-PV, evaluated ropeg against HU in patients with PV. PROUD-PV was powered to establish the non-inferiority of ropeg against HU with a composite primary endpoint of complete hematological response (CHR) and resolution of splenomegaly at 12 months; CHR and symptomatic improvement were the coprimary endpoints in the CONTINUATION-PV study. At a median follow-up of 182 weeks in the ropeg arm, 21%
(28% in HU arm at a median follow-up of 164 weeks) and
53% of patients (38% in HU arm, p = 0.044) met the pri-
mary endpoints in PROUD-PV and CONTINUATION-
PV study, respectively. CHR without the spleen criterion
in the ropeg arm was met in 43% (46% in HU, p = 0.63 at
12 months) and 71% (51% in HU, p = 0.012 at 36 months)
in the PROUD-PV and CONTINUATION-PV, respec-
tively. Liver enzyme abnormalities were the most fre-
quently reported grade 3/4 adverse events in the ropeg
arm and expected myelosuppression in the HU arm with
comparable rates between the groups. Neuropsychiatric
manifestations in the ropeg arm were rare [31]. Given
these encouraging results, Ropeg is currently approved
in Europe as a first-line agent for the treatment of PV in
the absence of symptomatic splenomegaly and is under
review for FDA approval in the USA.

Givinostat, a histone-deacetylase (HDAC) inhibitor has
demonstrated preclinical activity in selective targeting
of the JAK2 V617F clone by inhibiting the downstream
signaling [32]. Subsequently, several studies have shown
that givinostat is clinically active either as monotherapy
or in combination with HU [33, 34]. Most recently, givi-
nostat was evaluated in a dose-finding/proof of concept
study in patients with PV. Givinostat exhibited on-target
activity, and 100 mg twice daily was deemed the recom-
ended phase 2 dose (RP2D). In part B, proof of concept
phase, the ORR rate was 80.6% at the end of three cycles
and 50% of patients reported symptomatic improvement
(pruritus, headache) with givinostat treatment. Almost
all patients experienced grade 1/2 treatment-related
adverse event (TEAE) [diarrhea—51%; thrombocyto-
enia—45%; increased serum creatinine—37%]. Based
on these results, a registration trial of givinostat in PV
patients is underway [35].

The P53-MDM2 axis is a novel therapeutic target in
MPNs. MDM2 negatively regulates p53, promotes its
degradation as well as inhibits p53 transcription. Pre-
clinical studies have shown that MDM2 is upregulated
in JAK2 V617F-positive MPN hematopoietic progenitor
cells, resulting in low p53 RNA levels that has led to the
evaluation of MDM2 inhibitors in MPNs [36]. A recently
published proof of concept study of Idasanutlin, an oral
MDM2 inhibitor, in the second-line setting in patients
with high-risk PV/ET demonstrated an overall response
rate (ORR) of 58% (7/12) and a durable response
(16.8 months) with monotherapy. Idasanutlin was well
tolerated with no dose-limiting toxicities; low-grade
gastrointestinal toxicity (diarrhea/nausea in 80%) was
common but manageable with a scheduled antiemetic
regimen. Collectively, idasanutlin demonstrated safety
and on-target clinical activity in JAK inhibitor-naïve,
HU/IFN-resistant, or intolerant PV/ET patients. A global
phase II trial in HU refractory PV is underway [37].

(Lysine-specific demethylase 1 (LSD1) is an epigenetic
enzyme that maintains steady-state hematopoiesis and
LSD1 inhibition-abrogated erythropoiesis, granulopoie-
sis, and thrombopoiesis in a reversible fashion. Addition-
ally, LSD1 is found to be overexpressed in MPNs [38].
IMG7289 (bomedemstat), an LSD1 inhibitor, reduced
splenomegaly, normalized blood counts, and pro-
longed survival in the Jak2 V617F murine model [39],
which has led to the clinical evaluation of bomedemstat
as a second-line agent in PV and ET (NCT04254978)
(NCT04262141).

Furthermore, the recent understanding of the mecha-
nistic basis of CALR mutated MPN has revealed sev-
eral potential novel therapeutic targets, especially in
harnessing host immunity. CALR mutations generate a
novel positively charged C terminus in the CALR pro-
tein, which could be exploited as a potential shared neo-
antigen, as the physical interaction between CALR and
MPL is essential for CALR-induced myeloproliferation
[40, 41]. Additionally, studies have shown that CALR is
immunogenic and immune escape occurs in patients
with CALR-mutated MPN [42]. In this regard, CALR-
specific CD4+ T-cell clone, which demonstrated specific
cytotoxicity against autologous CALR-mutant cells, has
been generated [43], and these results have formed the
basis of a phase 1 CALR exon 9 peptide vaccine in CALR-
mutated MPNs. (NCT03566446) Most recently, Bozkus
et al demonstrated that a subset of patients with CALR-
mutated MPN exhibits specific T-cell responses against
the CALR C-terminus that is completely abrogated by the
expression of PD-1 or CTLA4. Ex vivo treatment with
an anti–PD-1 antibody restored mutant CALR-specific
T-cell responses in the peripheral blood mononuclear
cells of CALR-mutated MPN patients [44]. Clinical eval-
uation of a vaccine-based approach in combination with
a PD-1 inhibitor is underway.

Agents in clinical development in MF (Fig. 1)
Ruxolitinib, a JAK1/JAK2 inhibitor (2011) [45] and fed-
ratinib, a JAK2/FLT3 inhibitor (2019) [46] are approved
in the USA for MF patients with splenomegaly and/or
constitutional symptoms regardless of the presence
of mutated JAK2. Although long-term follow-up stud-
ies have validated the sustained benefit of ruxolitinib in
MF patients in terms of improvement in splenomegaly,
symptom burden, and quality of life with an increase
in overall survival (OS), a subset of patients are intoler-
ant or refractory to JAK inhibitor therapy. While the
median OS in ruxolitinib-treated patients is 60 months,
the median OS post ruxolitinib discontinuation drops significantly (14 months) [47, 48]. Furthermore, clonal evolution or the finding of platelets < 100 × 10^9/L at the time of ruxolitinib discontinuation was found to be associated with particularly poor prognosis in patients with MF. Additionally, Kuykendall et al. evaluated the clinical outcomes and salvage treatment options in patients who received and discontinued ruxolitinib. In 64 evaluable patients, new cytopenias (anemia—33%; thrombocytopenia—11%) were the most common reasons for an impediment to ruxolitinib continuation after a median treatment time of 3.8 months. Of note, 26% of patients responded to salvage treatment options leading to better outcomes than those who did not receive additional therapy, suggesting that responses were salvageable in some patients even after ruxolitinib discontinuation. However, these responses were rare, representing an area of unmet clinical need in ruxolitinib pretreated patients with MF [49]. Therefore, there is a constant drive to improve upon the existing treatment options in patients with MF. Currently, many novel therapeutic agents are in clinical development in the front-line setting (monotherapy), “Add on” with ruxolitinib as a complementary therapeutic strategy, second-line setting, or treatment directed at mitigation of cytopenias (Fig. 2).

**Front-line setting**

Pacritinib is a multikinase inhibitor of JAK2, FLT3, IRAK1, and CSF1R, with less myelosuppressive effect noted in the early-phase trials and further evidenced by the anemia response [25% achieved transfusion independence (TII)] and platelet improvement (35% increase in mean platelet count noted in those with a baseline platelet count lower than 50 × 10^9/L) in the PERSIST-1
randomized controlled trial in JAK inhibitor naïve patients with MF [50]. PERSIST-2, a randomized controlled trial, evaluated pacritinib in MF patients with either disease or therapy-related (ruxolitinib) thrombocytopenia (platelets < $10^9$/L), and they were randomized to two doses of pacritinib (200 mg BID or 400 mg once daily) or BAT (best available therapy), which could include ruxolitinib as well. Eighteen percent of patients enrolled on the pacritinib arms achieved ≥35% spleen volume reduction (SVR35%) compared with 3% on the BAT arm ($p=0.001$), and these improvements were more noticeable in the pacritinib 200 mg BID arm [≥35% SVR: 22% vs. 3%; $p=0.001$; ≥50% reduction in myelofibrosis-related total symptom score (TSS50): 32% vs. 14%; $p=0.01$]. Grade 3 or more thrombocytopenia, cardiac AEs, and therapy discontinuation were less frequent in the twice-daily arm [51]. Pacritinib development was interrupted due to the full clinical hold placed by the FDA in February 2016 due to safety concerns (increased hemorrhagic risk and mortality), which prompted an independent review that deemed mortality rates were not different between the study arms. Recently presented phase II PAC203 (NCT03165734) dose-finding (100 mg daily, 100 mg twice daily, and 200 mg twice daily) study evaluated pacritinib with preplanned built-in safety protocols for mitigating cardiac and hemorrhagic risk (concomitant anticoagulant/antiplatelet and QT-prolonging agents were contraindicated). Pacritinib was well tolerated, and 17% of patients with severe thrombocytopenia (<$50 \times 10^9$/L) attained spleen responses in the 200 mg BID cohort [52]. Given that thrombocytopenia (especially platelet count < $50 \times 10^9$/L) is a poor prognostic factor in patients with MF and ruxolitinib is only approved for those with a minimum platelet count of $50 \times 10^9$/L, pacritinib can potentially bridge this chasm and offer a viable therapeutic option for this challenging population subset. The PACIFICA phase III registration trial will evaluate the safety and efficacy of 200 mg BID of pacritinib compared to the physician's choice (low-dose ruxolitinib, corticosteroids, hydroxyurea, or danazol), in patients with MF and severe thrombocytopenia (<$50 \times 10^9$/L) and less than 12 weeks of prior JAK inhibitor therapy [53] (NCT03165734).

**Ruxolitinib “Add-on” strategies**

Itacitinib is a selective JAK1 inhibitor being evaluated in MF under the premise that selective JAK1 inhibition will abrogate proinflammatory signaling without affecting the JAK2-mediated hematopoiesis. A phase II open-label study evaluated the safety and efficacy of three dose levels [100 (n=10) or 200 mg BID [45], 600 mg QD [32]] of itacitinib in MF patients with TSS50 at week 12 as the primary endpoint. In total, 35.7% and 35.5% achieved the primary endpoint in the 200 mg BID and 600 mg QD as compared to 20% in the 100 mg BID cohort. Modest SVR was observed in the higher dose cohorts. Notably,
53.8% experienced a ≥ 50% reduction in the number of red blood cell units transfused, and fatigue was the most common TEAE [54]. Itacitinib is currently being evaluated in two cohorts with one cohort in combination with ruxolitinib and the other in JAK inhibitor failure/intolerance in patients with MF (NCT03144687).

Masarova et al. investigated the sequential combination of ruxolitinib with azacitidine, a hypomethylating agent, preceded by an initial run-in phase with ruxolitinib monotherapy. In total, 72% (33/46) of patients achieved an objective response per International Working Group for Myelofibrosis Research and Treatment (IWG-MRT) criteria with a median time to response of 1.8 months, and most responses occurred on ruxolitinib monotherapy. In total, 57% of patients experienced ≥ grade 1 improvement in bone marrow fibrosis (BMF). Of note, 20% (3/15) of patients with cytogenetic abnormalities at diagnosis achieved a complete cytogenetic response at the end of 12 months of combination therapy. The combination was relatively well tolerated with only transient grade 3/4 myelosuppression that did not warrant treatment interruption [55] (NCT01787487).

CPI-0610 is a novel Bromodomain and Extraterminal domain Protein (BET) inhibitor that is currently undergoing clinical evaluation with two treatment arms investigating the “add-on” approach with ruxolitinib in MF patients in the JAK inhibitor naïve [Arm(A)3] and experienced (A2) with suboptimal response settings and the other evaluating CPI-0610 monotherapy in patients after ruxolitinib discontinuation (A1). In the first-line setting (A3), 73% achieved SVR35% at 12 weeks; 59% achieved TSS50, and 46% experienced at least one-grade improvement in BMF. In the JAK inhibitor “experienced” cohort, patients were stratified by red blood cell transfusion status. While 34% of evaluable transfusion-dependent (TD) patients converted to TI in A2 (NCT02158858), 21% of TD patients converted to TI in A1 and SVR35% was comparable in A1 (24%) and A2 (22%). CPI-0610 was tolerable with minimal grade 3/4 myelosuppression, and thrombocytopenia, low-grade nausea, and vomiting were the most commonly observed TEAE [56, 57]. Given these encouraging results (spleen and symptom response) and the potential disease-modifying activity (improvement in anemia and bone marrow fibrosis), a phase III, double-blind, randomized study comparing combination CPI-0610 and ruxolitinib to ruxolitinib monotherapy will start in the fourth quarter of 2020 (MANIFEST-2).

Phosphatidylinositol 3-Kinase (PI3K) inhibitors are being evaluated in combination with ruxolitinib to improve upon the suboptimal response to ruxolitinib in MF patients. In a phase II study of combination umbralisib and ruxolitinib therapy in MF, 9% (2/23) of treated patients achieved a complete response (CR), and 48% (11/23) experienced clinical improvement. One patient experienced colitis, but other class-specific side effects (hepatotoxicity, pneumonitis) were not observed [58]. The recently presented interim study results of pasclisib, a potent and highly selective next-generation PI3Kδ inhibitor in combination with ruxolitinib. Pasclisib was evaluated in two dosing schedules (QD for eight weeks followed by weekly; daily). Recently presented data showed that the intensive daily dosing schedule was found to be more efficacious than the weekly schedule (median percent change in spleen volume: −13% vs. −2.3%; TSS: −51.4% vs. −14.0%, respectively, at week 12]. Pasclisib was well tolerated with no TEAE inherent to PI3K inhibitors (pneumonitis, colitis, diarrhea). Daily pasclisib “add-on” to ruxolitinib will be evaluated further in a planned phase 3 trial (NCT02718300) [59].

Navitoclax, a non-selective Bcl2 inhibitor, is being evaluated in combination with ruxolitinib to improve response in patients with MF. These patients were heavily pretreated (>3 lines of prior therapy), and 50% of treated patients harbored high molecular risk mutations (n=34). Thirty percent of evaluable patients achieved SVR35%, 35% achieved TSS50, and ≥1-grade BMF reduction was seen in 25% of patients suggesting disease-modifying activity. On target thrombocytopenia was the most common TEAE, but there were no grade ≥3 bleeding events or treatment-related deaths. The combination was well-tolerated, and this combination will be evaluated in randomized phase 3 trials in both treatment-naïve and JAK inhibitor-treated patients [60] (NCT03222609).

A phase I/II RUXOPEG adaptive design trial is evaluating the combination of ruxolitinib and pegylated interferon alpha-2a in treatment-naïve DIPSS intermediate- or high-risk MF patients on the basis that this combination may permit administration of lower doses of interferon and improve tolerability. Phase I will test different combinations of three dose levels of each drug, and phase II will randomize the two best dose combinations from the phase I. The primary endpoint is composed of safety and efficacy objectives as denoted by the dose-limiting toxicity (DLT) within 45 days and SVR50% in 24 weeks, respectively. Thus far, fifteen patients have been enrolled in phase I; no DLT has been observed in the highest dose tested (ruxolitinib 15 mg BID + IFNa 135 mcg/week), and an early signal for efficacy has been reported (three-partial responses, seven-hematological improvement). The trial is ongoing (NCT02742324) [61].

Second line: JAK inhibitor relapsed/ refractory/intolerant setting

The 5-year follow-up of the COMFORT-1 trial reported a median duration of approximately 3.2 years of splen...
response, suggesting that the disease response to JAK inhibitors is not everlasting [62]. Although progressive disease per IWG-MRT includes only new/progressive splenomegaly and increasing blast counts either in the blood or marrow, clinically patients may exhibit evidence of disease progression through worsening cytopenias or loss of symptom response [63]. Currently, the widely accepted definition of ruxolitinib failure is centered around spleen size and presence of cytopenias (Table 1) [52]. Furthermore, a retrospective claims database study reported that the median treatment progression-free survival after ruxolitinib discontinuation is six months, 95% (CI: 4.4, 8.3 months), which, coupled with poor outcomes post discontinuation, reiterates the urgent need to explore novel therapeutic options in MF patients experiencing ruxolitinib failure (relapsed/refractory or intolerant to ruxolitinib treatment) [64].

Momelotinib is a JAK1/2 inhibitor as well as a type 1 activin receptor (ACVR1) inhibitor being evaluated in MF patients with anemia on the premise that ACVR1 inhibition regulates hepcidin levels to restore iron homeostasis and improve anemia [66]. SIMPLIFY-1 study compared momelotinib with ruxolitinib in treatment-naive MF patients. Although the trial met the non-inferiority primary endpoint for ≥35% SVR at 24 weeks (26.5% for momelotinib versus 29% for ruxolitinib, \( p = 0.011 \)), it failed to meet the TSS50 endpoint. Notably, the momelotinib treatment arm enjoyed a higher rate of TI at week 24 than the ruxolitinib arm (66.5% vs. 49.3%, nominal \( p < 0.001 \)) [67]. SIMPLIFY-2 compared momelotinib to BAT (including ruxolitinib) in MF patients intolerant to ruxolitinib. The study failed to meet its primary endpoint SVR35%, but the TSS50 endpoint was met. Akin to SIMPLIFY-1, more momelotinib-treated patients achieved TI (43% vs. 21% nominal \( p = 0.0012 \)) [68]. However, in both trials, the hierarchal study design precluded the investigators from claiming the statistically significant anemia-related endpoints. Most recently, an open-label phase II study evaluated momelotinib in RBC TD patients with MF, and 34% achieved TI at week 24 [69]. The MOMENTUM trial will compare momelotinib to danazol in symptomatic and anemic patients with MF in the second-line setting (NCT04173494).

PRM-151 is a recombinant form of pentraxin-2, an endogenous serum amyloid protein that downregulates activated fibrogenic monocyte-macrophages activity in several organ models of fibrosis, including the bone marrow [70]. The first stage of phase II, open-label, extension study showed that PRM-151 was well tolerated as a monthly infusion either alone or in combination with ruxolitinib, and no unexpected AEs were observed in patients with MF. TSS50 was similar between both arms, and 44% of treated patients experienced at least a 1-grade reduction from grade 3 BMF at baseline (NCT01981850) [71]. In stage two, randomized, double-blind evaluation of three dose levels of PRM-151 infusional monotherapy in MF patients intolerant/refractory to JAK inhibitors, the primary endpoint was to determine the effective dose inducing at least a 1-grade reduction in BMF. All tested dose levels demonstrated greater than 1-grade BMF reduction, and the effect was similar across the tested doses [0.3 mg/kg: 30%; 3 mg/kg: 28%, and 10 mg/kg: 25%]. SVR35% was observed in only one patient. PRM-151 was well tolerated, and non-hematological AEs included fatigue, cough, and weight loss. Despite these encouraging findings, the further development of this drug in MF is uncertain as it evinced mostly BMF reduction. PRM-151 is currently undergoing registration trials in idiopathic pulmonary fibrosis [72].

Bomedemstat inhibits LSD1, an epigenetic target of interest in MPNs. LSD1 is essential for normal megakaryocyte function, and thrombocytopenia would be an expected dose-limiting side effect of LSD1 inhibition. In a phase II trial of bodememstat in the second-line setting (\( n = 31 \)), 12.5% of treated patients achieved SVR35%, 44.4% experienced TSS50, and ≥1-grade BMF reduction was noted in 15% of treated patients. Given the expectant thrombocytopenia, the dose up-titration of bodememstat was individualized to achieve a target platelet count of 50 × 10^9/L. No new safety signals or DLTs were observed. Further evaluation is underway (NCT03136185) [73].

Harnessing the targets in the apoptotic machinery has long been an object of clinical interest in MF. KRT-232,

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**Table 1 Criteria for ruxolitinib failure in patients with MF—adapted from [52, 65]**

| Criteria       | Ruxolitinib duration | Cytopenas                         | Spleen size                                                                 |
|----------------|----------------------|-----------------------------------|-------------------------------------------------------------------------------|
| Relapsed       | ≥ 3 months           | –                                 | Regrowth < 10% SVR or < 30% decrease in spleen size by palpation from baseline following an initial response |
| Refractory     | ≥ 3 months           | –                                 | < 10% SVR or < 30% decrease in spleen size by palpation from baseline         |
| Intolerant     | ≥ 28 days            | New grade ≥ 3 thrombocytopenia, anemia, hematoma/ hemorrhage or RBC transfusion requirement ≥ 2 units/month for 2 months | –                                                                            |

SVR spleen volume reduction, RBC red blood cell
a potent oral MDM2 inhibitor, is currently under clinical investigation in the second-line setting in patients with advanced MF. This study excludes patients who are intolerant to ruxolitinib and does not require a ruxolitinib washout period [74]. Patients were randomly assigned to either of the three-dose arms (120 (A1) or 240 mg (A3) daily for seven days in a 21-day cycle or 240 mg daily (A3) for seven days in a 28-day cycle). As 16% of patients achieved SVR35% in A3, 240 mg daily for seven days in a 28-day cycle is deemed to be the RP2D for further evaluation. In total, 51% of treated patients experienced grade 3 TEAE with gastrointestinal symptoms being the most common AEs [diarrhea (62%) and nausea (38%)] [75] (NCT03662126). KRT-232 is now being evaluated as combination therapy with ruxolitinib in a phase 1/2 trial enrolling patients with suboptimal response to single agent ruxolitinib (NCT04485260).

LCL-161 is an oral second mitochondrial activator of caspases (SMAC) mimetic that inhibits apoptosis and is administered on a weekly basis. The phase 2 study included all comers with advanced MF in the second-line setting with no restrictions for platelet count or previous HCT. Among 47 evaluable patients, 32% ORR was observed with most response improvement in symptom burden and anemia; only one patient had a spleen response. Fatigue was the most common cause for dose reduction, and low-grade nausea/vomiting was observed in 60% of the patients [76] (NCT02098161).

Alisertib is an aurora kinase A (AURKA) inhibitor that promotes megakaryocyte differentiation in MF and may mitigate bone marrow fibrosis. Alisertib was evaluated in patients with advanced MF in the second-line setting with a minimum platelet count ≥ 50 × 10^9/L and absolute neutrophil count ≥ 1 × 10^9/L. Alisertib was well tolerated, and spleen and symptom improvement were observed in 29% and 32% of patients, respectively. Most importantly, alisertib normalized the atypical morphology of megakaryocytes (restored the multilobed nuclei and abrogated clustering), and among the seven patients with available sequential marrow samples, five patients experienced >1-grade BMF, which correlated with the clinical responses (NCT02530619). The future development pathway for Alisertib is unclear [77].

Tagraxofusp is a CD123-targeted agent and currently approved in the treatment of blastic plasmacytoid dendritic cell neoplasm [78]. The shared phylogeny of plasmacytoid dendritic cells and monocytes, coupled with poor outcomes in MF patients with peripheral blood monocytosis, prompted the evaluation of tagraxofusp in the second-line setting. The study included all comers with no limitation in minimal platelet count at enrollment, and 26% of patients had documented monocytosis at baseline. Tagraxofusp was administered intravenously for three consecutive days in a 28-day cycle. Among 20 evaluable patients, 35% experienced objective clinical improvement, and 53% with baseline splenomegaly had some degree of reduction in spleen size as their best response. Tagraxofusp was reasonably well tolerated, with one patient experiencing grade 3 capillary leak syndrome [79] (NCT02268253).

Imetelstat is a competitive inhibitor of the telomerase enzyme complex comprising the RNA template with reverse transcriptase activity (hTERT). In a proof of concept study of 33 patients with advanced MF, imetelstat evinced an ORR of 21% limited to those with JAK2, SF3BI, or U2AF1 mutations. The study did not show on-target activity (telomerase length) [80]. The subsequent phase 2, global IMBARK trial evaluated two dose levels of imetelstat (4.7 mg/kg and 9.4 mg/kg) administered intravenously every three weeks in 107 patients with advanced MF in the second-line setting (NCT02426086). Although SVR35% (10%) and TSS50 (32%) were only modest in the higher dose arm, the median survival was 29.9 months as compared with 19 months in the low-dose arm and the reported median survival of 13–14 months following ruxolitinib discontinuation [81]. Furthermore, imetelstat exhibited on-target activity and brought about greater than 50% reduction of hTERT expression levels, which correlated with clinical responses and longer OS in the 9.4 mg/kg arm [82]. Most importantly, the survival advantage of imetelstat was validated in a real-world cohort using a closely matched propensity score analysis [30.69 mo (95% CI 25.2, not estimable) vs. 12.04 mo (95% CI 9.5, 16.6) (BAT)] [83]. Given these encouraging results, the phase III registration trial of imetelstat is soon underway with OS as the primary endpoint, a novel endpoint that has never been explored in the drug development landscape of MF.

**Drugs targeting cytopenias in MF**

Anemia independently predicts shortened survival in MF, and TD-anemia categorizes an MF patient in the higher-risk category regardless of the presence or absence of other adverse risk factors [84]. Furthermore, anemia is the most common reason for ruxolitinib discontinuation [49]. Several drugs are in clinical development for mitigating anemia in MF so as to safely continue MF-directed therapy. MPNSG-0212, a German study, evaluated pomalidomide in combination with ruxolitinib [two dose levels of pomalidomide: fixed low dose (A1) and dose escalation up to 2 mg (A2)] in MF patients with anemia ≤ RBC-TD. The A1 cohort exhibited an ORR of 18%, and there was a trend to sustained hemoglobin improvement with longer durations of treatment. TEAE was comparable between both arms with pneumonia and sepsis being the most grade ≥ 3 AEs [85].
Sotatercept and luspatercept are erythroid maturation agents (EMA) that act as activin receptor ligand traps of IIA and IIB, respectively [86, 87]. They are administered subcutaneously every three weeks, and luspatercept is currently approved for the treatment of anemia in low-risk myelodysplastic syndrome with ringed sideroblasts [88]. Sustained hemoglobin increase ≥1.5 g/dL for ≥12 consecutive weeks in TI patients or achieving RBC-TI in TD patients is the primary endpoints in the clinical trial evaluation of these EMAs in MF. Sotatercept
monotherapy demonstrated an ORR of 35%, of which three patients achieved RBC-TI [89] (NCT01712308). In the recently presented study of luspatercept mono-
therapy and combination therapy with ruxolitinib in ane-
mic patients with MF, 10% of treated patients each in the
luspatercept monotherapy arm and 21% and 32% in the
combination therapy arm achieved the primary endpoint
in TI and TD patients, respectively (NCT03194542) [90].
Hypertension and bone pain were the most common,
class-specific TEAE shared by both drugs. Further evalu-
ation is ongoing, and a phase 3 trial is being planned.

Disease-related thrombocytopenia is an adverse prog-
nostic factor in MF, which often precludes these patients
from treatment with a JAK inhibitor or leads to dose
attenuation resulting in suboptimal responses. Thalido-
mide in combination with prednisone has evoked mod-
est improvement in platelet counts in patients with MF
[91]. Most recently, a study of low-dose thalidomide in
combination with ruxolitinib in patients with MF in the
second-line setting (relapsed/refractory) showed an ORR
of 60%, and platelet response was observed in 75% of
patients with baseline thrombocytopenia. The combina-
tion was well tolerated with one patient each experienci-
ing a thromboembolic event and grade 3 neutropenia.
This combination may allow for optimal dosing of rux-
olitinib in MF patients with baseline thrombocytopenia
[92].

Several other agents exploiting the interconnected
pathological pathways in MF are in various stages of
early-phase clinical development (Table 2).

Conclusion
Advances in diagnostic techniques, i.e., next-genera-
tion sequencing, single-cell transcriptome approaches,
have carefully refined the molecular signature of MPNs,
leading to enhanced insight on clonal dynamics and
architecture, thereby informing rationally based treat-
ment approaches. Although HU or IFNα is the front-
line agent in the treatment of PV, 25% of patients are
intolerant to these agents and experience disease pro-
gression while receiving therapy. In light of this, ongo-
ing translational research endeavors have identified
mechanistic-based targeted therapeutic agents that
may improve the outcomes in PV. Comparably in MF,
sustained disease-modifying activity or durable remis-
sions are not seen with the currently approved JAK
inhibitors, i.e., ruxolitinib and fedratinib. Therefore, it
is crucial to improve upon the existing understanding
of the disease and treatment-resistant mechanisms in
MF. As such, research efforts are ongoing to develop
novel JAK inhibitors or drugs with distinct mechanisms
of action that offer a better side effect profile and toler-
ability in patients with MPNs. Ropeginterferon in low-
risk PV, pacritinib in the front-line setting of extreme
thrombocytopenia, CPI-0610 combination therapy in
JAK inhibitor-naïve patients, imetelstat in the second-
line setting to improve survival outcomes, and luspa-
tercept for the treatment of MF patients with anemia
are some of the promising agents that look to achieve
results in phase 3 trials and gain regulatory approval for
the treatment of MPNs.

Abbreviations
JAK: Janus-associated kinase; STAT: Signal transducers and activators of trans-
scription; MPN: Myeloproliferative neoplasms; PV: Polycythemia vera; ET: Essential
thrombocythemia; PMF: Primary myelofibrosis; USFDA: The United States
Food and Drug Administration; HCT: Hematopoietic cell transplantation; HU: Hy-
donxyurea; IFNα: Pegylated interferon α; Ropeg: Ropeginterferon alfa-2b;
MPD-RC: Myeloproliferative disorders research consortium; CHR: Complete
hematological response; HDAC: Histone-deacetylase; TEAE: Treatment-related
adverse event; LSD1: Lysine-specific demethylase 1; TI: Transfusion independ-
ence; TD: Transfusion dependence; SVR35%: ≥ 35% Spleen volume reduction;
ING-MRT: International Working Group for Myelofibrosis Research and
Treatment; BAT: Best available therapy; BMF: Bone marrow fibrosis; TSS50: ≥
50% Reduction in myelofibrosis-related total symptom score; A: Arm; BET:
Bromodomain and Extraterminal domain Protein; PI3K: Phosphatidylinositol
3-Kinase; DLT: Dose-limiting toxicity; ACV: Activin receptor; SMAC: Second
mitochondrial activator of caspases; AURKA: Aurora kinase A; TERT: RNA tem-
plate with reverse transcriptase.

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