Ruxolitinib: a targeted treatment option for patients with polycythemia vera

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Abstract: Polycythemia vera (PV) is a chronic myeloproliferative neoplasm characterized by erythrocytosis and the presence of Janus kinase (JAK) 2V617F or similar mutations. This review summarizes the pathophysiology of PV, the challenges associated with traditional treatment options, and the scientific rationale and supportive clinical evidence for targeted therapy with ruxolitinib. Accumulating evidence indicates that activating mutations in JAK2 drive the PV disease state. Traditional PV treatment strategies, including aspirin, phlebotomy, and cytoreductive agents such as hydroxyurea, provide clinical benefits for some but not all patients and may not adequately treat PV-related symptoms. Furthermore, traditional treatment approaches are associated with potential side effects that may limit their usage and lead some patients to discontinue the treatment. Ruxolitinib is an orally available small-molecule tyrosine kinase inhibitor that is a potent and selective inhibitor of JAK1/JAK2. Ruxolitinib is approved in the US for patients with PV with an inadequate response or intolerance to hydroxyurea and in Europe for adults with PV who are resistant to or intolerant of hydroxyurea. In the Phase III RESPONSE registration trial, ruxolitinib was superior to the best available therapy in patients with PV who were resistant to or intolerant of hydroxyurea in controlling hematocrit levels, reducing spleen volume, and improving PV-related symptoms and quality-of-life measures. The most common nonhematologic adverse events in ruxolitinib-treated patients were headache, diarrhea, pruritus, and fatigue in the RESPONSE trial; hematologic adverse events were primarily grade 1 or 2. In the Phase IIIb nonregistration RELIEF trial, there were nonsignificant trends toward an improved symptom control in patients with PV on a stable hydroxyurea dose who were generally well controlled but reported disease-associated symptoms and switched to ruxolitinib vs those who continued hydroxyurea therapy. Updated treatment guidelines will be important for educating physicians about the role of ruxolitinib in the treatment of patients with PV.

Keywords: myeloproliferative disorder, polycythemia vera, Janus kinase inhibitor

Introduction
Polycythemia vera (PV) is a disorder predominantly characterized by erythrocytosis.1 As opposed to secondary erythrocytoses, PV and primary familial congenital polycythemia are categorized as primary erythrocytoses, which result from enhanced responses to erythropoietin (EPO).2 In 1951, PV and three other disorders with similar pathophysiologic characteristics (myelofibrosis [MF], essential thrombocythemia [ET], and chronic myeloid leukemia) were characterized as “myeloproliferative disorders” by Dr William Dameshek.3 Subsequent cytogenetic analyses and clonality studies demonstrated that PV is a clonal malignancy acquired through one or more somatic mutations in a pluripotent hematopoietic stem cell, leading to increased myelo...
proliferation. However, a molecular target responsible for PV was not yet identified at that time.

Early hypotheses regarding the pathologic basis of PV included potential abnormal growth factor signaling pathways, transcriptional dysregulation, and constitutive activation of signal transducer and activator of transcription (STAT) proteins. Polycythemia rubra vera-1 (PRV-1) was once hypothesized as a PV-specific marker that could be important for elucidating the molecular mechanism of the disorder. The PV diagnostic strategy drastically changed after the discovery of activating mutations in the JAK2 tyrosine kinase. Following this discovery, the World Health Organization characterized PV as a chronic myeloproliferative neoplasm (MPN) primarily defined by erythrocytosis and the presence of JAK2V617F or similar mutations.

Patients diagnosed with PV may have a marked disease burden and a higher mortality risk when compared with the general population, primarily driven by cardiovascular/thrombotic events, disease transformation to MF or acute myeloid leukemia (AML), and solid malignancies. Traditional PV treatment strategies include aspirin, phlebotomy, hydroxyurea, and other cytoreductive treatments. Such options provide clinical benefits for some patients, including a reduced risk of cardiovascular/thrombotic events. However, traditional treatment options do not provide adequate benefit for some patients and may not alleviate PV-related symptoms. As such, there remains an unmet need for improved clinical outcomes, symptom alleviation, and enhancement of quality-of-life (QoL).

The objective of this review is to provide a summary of the pathobiology of PV, including the evolving understanding of the molecular etiology of PV, the challenges associated with traditional PV treatment options, and the scientific rationale and clinical data for ruxolitinib (Jakafi, Incyte Corporation, Wilmington, DE, USA; Jakavi, Novartis AG, Horsham, West Sussex, UK) supporting its role as a new targeted treatment option for patients with PV.

**Biology of PV: pre- and post-JAK2V617F era**

Pre-JAK2V617F molecular understanding of the etiology of PV

Early studies of erythropoiesis provided important information about the hematopoiesis process in the PV setting. In vitro analyses demonstrated that bone marrow progenitor cells isolated from patients with PV (but not control bone marrow samples) were able to form EPO-independent endogenous erythroid colonies (EECs). In 1987, a review of nine studies reported that 97% of patients with PV had EECs in the bone marrow and/or peripheral blood. Although EECs were also observed in some patients with ET, their presence is a hallmark of PV and was used as a clinical diagnostic tool.

The observed increased proliferative responsiveness of PV progenitor cells to EPO, IGF-1, and other growth factors (eg, interleukin [IL]-3, granulocyte-macrophage colony-stimulating factor [GM-CSF], thrombopoietin, and stem cell factor) implicated abnormal cytokine signaling pathways in the molecular underpinnings of PV. However, studies that examined mutations related to cytokine signaling targets (eg, the EPO receptor, IGF-1 receptor, IGF-1–binding proteins, and tyrosine phosphatases) were unsuccessful in elucidating the pathogenesis of PV. It was also hypothesized that abnormal cytokine signaling was not necessarily related to a limited number of specific mutations but rather to more general defects in transcriptional regulation that could affect a variety of metabolic pathways that play a role in the pathogenesis of PV. The transcriptional dysregulation hypothesis was supported by studies of cells from patients with PV, which reported decreased levels of the thrombopoietin receptor c-MPL and a proportion of erythroid progenitors expressing BCL-x (an antiapoptotic protein). Downstream signal transduction molecules important to cytokine receptor signaling, including EPO-mediated pathways, were further studied to identify the potential PV candidate genes. It was hypothesized that EPO-mediated activation of the JAK/STAT pathway induced BCL-x expression, which inhibited apoptosis. However, data suggested that constitutive STAT3 activation alone was not the primary molecular cause of PV.

Collectively, available data up to this point suggested that the JAK/STAT transduction pathway might play a critical role in preventing the apoptosis of erythroid progenitors but not necessarily in the proliferation of erythroid cells. The search for the molecular cause of PV continued. Using subtractive hybridization techniques, overexpression of PRV-1 mRNA was observed in the granulocytes of patients with PV but not in normal controls. However, the specificity of PRV-1 mRNA overexpression to the PV setting was later contradicted by data indicating no consistent differences in PRV-1 protein levels between granulocytes from patients with PV and those from normal controls.
Furthermore, the sensitivity of PRV-1 expression to GM-CSF exposure suggested that alterations in cytokine levels (not the PV disease state itself) explained the variability of PRV-1 levels. As such, the technically demanding and time-consuming EEC assay continued to be the most reliable test for the diagnosis of PV, and the molecular cause of PV remained elusive.

**Discovery of JAK2V617F and the importance of JAK2 in the pathogenesis of PV**

In 2005, four separate groups identified the somatic gain-of-function JAK2V617F mutation, a constitutively active allele associated with activation of downstream signaling components, including STAT3 and STAT5. In murine models, JAK2V617F bone marrow cells are associated with PV disease features, including erythrocytosis, leukocytosis, and enlarged spleen. Nearly all patients with PV have an activating mutation in JAK2, most often JAK2V617F (95% of patients) or an activating mutation in exon 12 (4%) of patients. Furthermore, patients with elevated JAK2 allele burden (percentage of mutant allele relative to the total [wild type + mutant]) are at increased risk of developing post-PV MF.

The discovery of the JAK2V617F mutation was critical for understanding the pathobiology of PV and dovetailed with other preclinical data concerning the JAK/STAT signaling pathway. For example, GM-CSF activates Jak1 and drives granulopoiesis in a pathway that includes JAK2 in cell culture systems, and IL-12–driven T-cell proliferation requires Jak2 activity. EPO signaling through the EPO receptor activates Jak2, which in turn activates STAT1, STAT3, and possibly STAT5 in cell culture systems. Mice lacking Jak2 are embryonic lethal because of extreme anemia. Unlike wild-type mice, hematopoietic progenitor cells from Jak2−/− mice do not proliferate and form megakaryocytic colonies in response to thrombopoietin. In addition, conditional knockout of Jak2 in adult hematopoietic progenitor cells is associated with reduced viability and lowered platelet counts. GM-CSF antiapoptotic activity in human eosinophils requires Jak2 activation. Finally, Jak2 is activated in response to thrombopoietin binding to the Mpl receptor and is required for the downstream activation of STAT3. With the knowledge that activating mutations in Jak2 were driving the PV disease state, researchers and clinicians had the rationale for the development of new, targeted treatment options, which was the impetus for testing ruxolitinib, a potent Jak1/Jak2 inhibitor, in patients with PV.

**Clinical presentation of PV**

**Elevated blood counts**

Elevated blood counts are one of the key diagnostic criteria associated with PV. In a large international study evaluating prognosis and survival among 1,545 patients with PV, many patients presented with elevated hemoglobin (median, 18.4 g/dL) and hematocrit values (median, 55%) at diagnosis. White blood cell (WBC) and platelet counts were also elevated (median, 10.4×10^9/L and 466×10^9/L, respectively). In addition, approximately one half of the patients presented with leukocytosis (WBC count >10.5×10^9/L; 49%) and thrombocytosis (platelet count ≥450×10^9/L; 53%).

Patients with PV who have a high JAK2 allele burden may experience further elevated blood counts compared with those who have less allele burden. A prospective study including 173 patients with PV demonstrated that there was a correlation between the JAK2V617F allele burden and the increased risk of erythropoiesis and myelopoiesis. Patients with higher JAK2V617F burden at diagnosis also presented with higher hematocrit levels (regression coefficient [β] =-0.67; P<0.001) and higher WBC counts (r=0.54; P<0.001).

**Thromboembolic and mortality risk**

Although elevated blood counts are important diagnostic markers of PV, thromboembolic events are often responsible for the initial clinical presentation of the disease. Patients with PV are at an increased risk of cardiovascular/thromboembolic events and mortality compared with the general population. Results from a large Swedish registry database study (n=11,155 patients with PV and n=44,620 matched controls) indicated that the risk of arterial and venous thrombosis in patients with PV was five- and ninefold higher, respectively, in patients with PV compared with the general population.

In a large, randomized, controlled trial testing cytoreductive therapy in patients with PV (CYTO-PV), elevated hematocrit levels of 45%–50% (n=183) were associated with a fourfold increase in cardiovascular complications and a significantly higher rate of cardiovascular death compared with those who had hematocrit level of <45% (n=182). Data from this trial also indicated that patients with a WBC count of ≥11×10^9/L were nearly four times more likely to experience major thrombosis when compared with those who had a WBC count of <7×10^9/L. However, available data do not suggest that thrombocytosis is a significant risk factor for cardiovascular/thromboembolic events in patients with PV.
the rates of thromboembolic events and mortality were not significantly different between patients with baseline platelet counts >400×10^9/L vs those with lower platelet counts. The molecular pathway responsible for the increased incidence of thromboembolic events in patients with PV is unclear. However, data suggest that constitutive JAK2 signaling may contribute to several PV features associated with the development of thromboembolic events, including erythrocytosis, increased adhesive qualities of JAK2V617F erythrocytes, leukocytosis, systemic inflammation, and activation of blood cell types.

**Signs and symptoms**

Patients with PV may experience a variety of disease-related symptoms, reduced QoL, and reduced work productivity. Fatigue, pruritus, difficulty in sleeping, day or night sweats, and dizziness are among the most frequently reported symptoms of PV. Patients with PV may also develop splenomegaly, which can be uncomfortable or painful in some patients.

The specific pathways underlying PV-related symptoms are unknown, but some evidence suggests that cytokine signaling plays an important role. Notably, serum levels of inflammatory cytokines, which signal through JAK1 and/or JAK2, are elevated in patients with PV. A cytokine profiling study in patients with MF found several clinical correlations between cytokine levels, constitutional symptoms, and splenomegaly.

Some symptoms may stem from increased blood counts caused by the constitutive activation of JAK2. The JAK2V617F mutation is associated with elevated hematocrit levels in murine models and in patients with PV, which may be associated with increased blood viscosity and symptoms that result from an impaired cerebral blood flow, including headache and dizziness. Patients with PV who have higher JAK2V617F allele burden may be at increased risk of developing splenomegaly and pruritus and are more frequently in need of cytoreductive therapy.

The effects of PV on symptoms and QoL, as commonly measured by the MPN Symptom Assessment Form (MPN-SAF) and the European Organisation for Research and Treatment of Cancer Quality-of-Life Questionnaire Core 30, are similar to those of other MPNs. Some patients report that burdensome PV-related symptoms have negative effects on their QoL, productivity, and activities of daily living. In a recent US survey of 380 patients with PV, 66% experienced symptoms that reduced their QoL, 22% were sick from work for ≥1 day in the preceding 30 days, and 48% experienced PV-related interference with daily activities.

**Limitations of traditional treatment options**

**Treatment overview**

Traditional treatment options are effective for many patients with PV but do not target the molecular underpinnings of the disease and are associated with several limitations, suggesting that improved treatment options are warranted. The goals of treatment in PV are to reduce the risk of cardiovascular/thrombotic events and control disease-related symptoms. Maintaining a hematocrit level of <45% is associated with a reduced risk of cardiovascular/thrombotic events and related death. For patients with splenomegaly, treatment should also aim to reduce spleen size. Aspirin, phlebotomy, and cytoreduction are the three main elements of traditional PV treatment strategies.

**Aspirin**

The European Collaboration on Low-Dose Aspirin in Polycythemia Vera clinical trial demonstrated that treatment with low-dose aspirin was associated with a reduced risk of cardiovascular/thrombotic events. Subsequently, a Cochrane meta-analysis reported that low-dose aspirin was associated with a nonsignificant reduction in the risk of fatal thrombotic events. However, patients receiving regular aspirin treatment (≥2×325 mg/wk) may be at a dose-dependent increased risk of gastrointestinal bleeding. Aspirin is contraindicated in patients with acquired von Willebrand syndrome and/or platelet count of >1,000×10^9/L, which are associated with an increased risk of bleeding.

**Phlebotomy**

Phlebotomy with or without cytoreductive treatment to maintain a hematocrit level of <45% was associated with reduced cardiovascular death and major thrombotic events in patients with PV. However, hematocrit maintenance at <45% with phlebotomy and other traditional treatment options can be difficult. The CYTO-PV study required hematocrit control within predefined limits with phlebotomy and/or platelet count of >1,000×10^9/L, which are associated with an increased risk of bleeding. Phlebotomy procedures may be poorly tolerated by some patients, as evidenced by the observation that 28% of patients in the high-hematocrit arm of the CYTO-PV trial discontinued phlebotomy treatment. In a large study of
phlebotomy patients undergoing venipuncture (N=3,315), 15% of patients reported that they feared the phlebotomy procedure and 3% reported that they avoided the procedure because of that fear. In addition, frequent phlebotomies can cause iron deficiency, which may be associated with restless leg syndrome and impairments in cognitive functioning and mental health in some patients.

**Cytoreductive agents**

Cytoreductive treatment with hydroxyurea is associated with increased survival and may be associated with reduced thromboembolic risk in patients with PV. However, one retrospective study indicated that 11% and 13% of patients treated with hydroxyurea become resistant and intolerant, respectively. In a multivariate analysis, resistance to hydroxyurea was associated with a 6.8-fold higher risk of transformation to AML or MF and a 5.6-fold higher risk of death. Hydroxyurea may also be associated with several hematologic and nonhematologic side effects that could limit treatment, including leg ulcers and other mucocutaneous manifestations, gastrointestinal toxicity, and fever.

Some data suggest that hydroxyurea may increase the risk of transformation to leukemia, especially in younger patients, although further analyses are required to confirm this hypothesis. The PV Study Group-08 trial reported that the incidence of AML at a median follow-up of 8.6 years was 5.9% in 51 hydroxyurea-treated patients compared with 1.5% in a historical control group of 134 patients from the pipobroman group. The authors acknowledged that the incidence of transformation to AML/MDS at 10 years, 15 years, and 20 years were 6.6%, 13.1%, 34.1%, and 52.1%, respectively, in the pipobroman group. The authors acknowledged that the incidence of transformation to AML/MDS in hydroxyurea-treated patients was higher than previously reported. However, they emphasized that the natural evolution of PV should be considered when interpreting the study results. The leukemogenic risk of hydroxyurea appears to be increased when treatment is preceded or followed by alkylating agents (eg, busulfan). Because hydroxyurea may modestly increase the risk of AML development, it has been suggested that this drug should be used with caution in younger patients (<40 years of age) with MPNs.

Interferon-α (IFN-α) and pegylated (PEG)-IFN-α variants are not currently indicated by the US Food and Drug Administration (FDA) or the European Medicines Agency for the treatment of patients with PV. However, IFN-α was recommended by the European LeukemiaNet in 2011 as second-line cytoreductive therapy for patients with PV who become resistant to or intolerant of hydroxyurea. Treatment with IFN-α and PEG-IFN-α variants has been associated with hematocrit control without phlebotomy, normalization of blood cell counts, and reductions in enlarged spleen size. In a review article that summarized clinical trial experience with different IFN-α variants in patients with PV published between 1991 and 2008, 60% (182/303) of patients achieved freedom from phlebotomies. An objective response was observed in ~80% of patients, although response criteria were heterogeneous among these studies.

In addition, some data suggest that IFN-α treatment may be associated with improvements in some PV-related signs and symptoms, including pruritus and splenomegaly. However, the route of administration (injection) and safety concerns about recombinant IFN-α formulations suggest that IFN-α may not be an ideal treatment option for some patients. A multicenter observational study reported that nonadherence to IFN-α primarily resulted from patients forgetting to administer the injection (50.2%) and other injection-related reasons (32.0%). Side effects of IFN-α, including influenza-like symptoms, autoimmune disorders, depression, cardiovascular disease, and ocular disease, may lead some patients to discontinue the treatment. These safety concerns are primarily associated with recombinant IFN-α variants, and it is important to note that PEG-IFN-α variants are associated with fewer toxicity-related treatment discontinuations. PEG-IFN-α2a has also demonstrated decreases in JAK2V617F allele burden over time and is associated with preferable hematologic responses in patients with PV.

In two Phase II clinical trials evaluating PEG-IFN-α in patients with PV, complete hematologic response was observed in 70% and 94.6% of patients (median follow-up time, 21 months and 31.4 months, respectively), and complete molecular response (undetectable JAK2V617F) was observed in 14% and 24.1% of evaluable patients, respectively. Discontinuations because of PEG-IFN-α-related adverse
Challenges of PV-related symptom alleviation

Traditional treatment options may not alleviate the PV-related symptom burden. In a survey-based study, patients with MPNs treated with hydroxyurea, IFN-α, busulfan, or reported similar MPN-SAF symptom severity scores as patients who did not receive myelosuppressive agents. Furthermore, a prospective analysis of patients with PV reported that treatment with hydroxyurea, aspirin, IFN-α, or phlebotomy was not associated with significant improvements in the MPN-SAF total symptom score (TSS). Collectively, available data suggest that a targeted treatment approach aimed at the molecular pathway associated with the pathogenesis of PV could be more effective than traditional treatment options for ameliorating PV-related symptoms.

Ruxolitinib: mode of action, pharmacodynamics, and pharmacokinetics

Mode of action and pharmacodynamics

Ruxolitinib phosphate (−(R)-3-(4-(7H-pyrrolo[2,3-d]pyrimidin-4-yl)-1H-pyrrozol-1-yl)-3-cyclopentylpropanenitrile phosphate) is a small-molecule tyrosine kinase inhibitor approved by the FDA for the treatment of patients with PV who have had an inadequate response to or are intolerant of hydroxyurea. In Europe, ruxolitinib is indicated for the treatment of adult patients with PV who are resistant to or intolerant of hydroxyurea. In the US, ruxolitinib is also indicated for the treatment of patients with intermediate- or high-risk MF, including primary MF, post-PV MF, and post-ET MF; ruxolitinib is approved for a similar population of patients with MF in Europe.

In preclinical models, ruxolitinib effectively inhibits the Jak/STAT signaling pathway as a potent and selective inhibitor of JAK1 (mean [SD] IC50 50%, 3.3 [1.2] nM) and JAK2 (mean [SD] IC50 2.8 [1.2] nM). Additionally, ruxolitinib has modest selectivity against Tyk2 (mean [SD] IC50 19 [3.2] nM) and marked selectivity against JAK3 (mean [SD] IC50 428 [243] nM). Ruxolitinib showed no significant inhibition against 26 additional kinases at concentrations 100-fold the IC50 of JAK1 and JAK2, suggesting high specificity for JAK1 and JAK2. The effectiveness of ruxolitinib in this pathway has also been demonstrated in cytokine-stimulated whole-blood assays in which preincubation with ruxolitinib inhibited IL-6– and thrombopoietin-mediated STAT3 phosphorylation (mean [SD] IC50 282 [54] nM and 281 [62] nM, respectively).

Several preclinical experiments demonstrated that ruxolitinib inhibits JAK pathway signaling of both wild-type and mutant JAK2. In cell lines expressing a JAK2V617F mutation, ruxolitinib effectively inhibited the phosphorylation of JAK2 downstream targets (eg, STAT3, STAT5, and ERK1/2) and induced apoptosis in a dose-dependent fashion. Ruxolitinib also inhibited erythroid and myeloid progenitor cell proliferation in primary cultures from healthy individuals (burst-forming unit-erythroid IC50 407 nM; colony-forming unit-erythroid IC50 551 nM) and patients with PV carrying the JAK2V617F mutation (burst-forming unit-erythroid IC50 223 nM; colony-forming unit-erythroid IC50 444 nM). In a murine model, ruxolitinib prolonged the survival, reduced the JAK2V617F allele burden, ameliorated splenomegaly, and normalized the elevated levels of circulating proinflammatory cytokines associated with debilitating constitutional symptoms of PV (eg, IL-6 and tumor necrosis factor-α).

Pharmacokinetics

Ruxolitinib pharmacokinetics supports an oral route of administration. Ruxolitinib has a dose-proportional systemic exposure profile, with minimal accumulation following multiple doses. In addition, ruxolitinib is rapidly absorbed, with 95% oral bioavailability. The maximum tolerated dose of ruxolitinib was established at 25 mg twice daily and 100 mg once daily, and the average terminal half-life was ~3 hours. Ruxolitinib is metabolized primarily by the cytochrome P450 3A4 (CYP3A4) enzyme. In healthy human subjects, >99% of ruxolitinib doses are metabolized in a pathway that includes oxidation to single and multiple hydroxylated products, some of which then undergo O-glucuronidation. Unmetabolized ruxolitinib is the predominant plasma drug entity until 6 hours postdose (58%). Most (96%) of the total dose is excreted within 24 hours postdose, primarily via urine (74%) and feces (22%). Patients who receive concomitant potent CYP3A4 inhibitors may require a 50% reduction in the ruxolitinib starting dose; however, data suggest that no dosage change is required when ruxolitinib is coadministered with inducers or mild to moderate inhibitors of CYP3A4.
Ruxolitinib dose reduction is recommended in patients with hepatic or renal impairment.\(^9\) In patients with mild and severe hepatic impairment, exposure to ruxolitinib (area under the curve) was increased because of the reduced clearance, although the magnitude of increase in ruxolitinib exposure was not correlated with the degree of hepatic impairment.\(^10\)

In contrast, an increasing severity of renal impairment was associated with increased exposure to ruxolitinib’s active metabolites and consequently with increased ruxolitinib pharmacologic activity.\(^10\)

**Ruxolitinib: clinical outcomes**

**Efficacy**

The randomized, open-label Phase III RESPONSE registration trial demonstrated that ruxolitinib was superior to the best available therapy in adult patients with PV who were phlebotomy dependent, had splenomegaly, and were resistant to or intolerant of hydroxyurea.\(^2\) Patients were randomized to receive ruxolitinib (n=110) or the best available therapy (n=112). Pharmacological treatment in the best available therapy was chosen at the discretion of the treating physician, primarily hydroxyurea (58.9%) and IFN (11.6%); 15.2% of patients received no medication (except aspirin).\(^3\) Several limitations of the RESPONSE trial, including the open-label design and the nonstandardized determination of treatment in the best available therapy arm, may have influenced the study treatment compliance and precluded the study from comparing specific treatments with ruxolitinib. However, the study was not powered for such comparisons, and allowing treating physicians to determine the course of treatment was representative of real-world clinical settings. It is not uncommon for patients to continue treatment with hydroxyurea despite the evidence of resistance or intolerance.

In the primary analysis, a higher proportion of patients receiving ruxolitinib vs the best available therapy achieved the composite primary endpoint of hematocrit control and \(\geq 35\%\) reduction in spleen volume from baseline at Week 32 (20.9% vs 0.9%, \(P<0.001\)).\(^2\) Higher proportions of patients receiving ruxolitinib also achieved individual components of the primary endpoint (hematocrit control, 60.0% vs 19.6%; \(\geq 35\%\) reduction in spleen volume from baseline, 38.2% vs 0.9%)\(^2\) and complete hematologic response (23.6% vs 8.9%, \(P=0.003\))\(^2\) at Week 32 (Table 1). A post hoc analysis demonstrated that the degree of splenomegaly at baseline did not influence the achievement of hematocrit control or spleen size reduction with ruxolitinib treatment.\(^10\) Data also suggest that ruxolitinib conferred benefits in patients who did not meet the primary study endpoint. In patients who did not achieve hematocrit control at Week 32, ruxolitinib prolonged the median time to subsequent phlebotomy eligibility compared with the best available therapy (52 weeks vs 21 weeks, respectively).\(^10\)

A total of 96 patients (85.7%) initially randomized to receive the best available therapy crossed over to the ruxolitinib arm because of lack of efficacy; most did so at or soon after the Week 32 visit.\(^2\) After cross over to ruxolitinib, a greater proportion of patients achieved \(\geq 35\%\) reduction in spleen size (crossover, 38.5%; best available therapy during randomized treatment, 1.8%), and phlebotomy treatment rates were lower (crossover, 38.5/100 patient-years; best available therapy during randomized treatment, 196.8/100 patient-years).\(^10\)

A preplanned analysis of RESPONSE conducted 80 weeks after the last patient received his/her first dose demonstrated a durable response with ruxolitinib. This analysis indicated that of the 23 patients randomized to ruxolitinib who achieved the primary endpoint response at Week 32,

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**Table 1** RESPONSE Phase III trial efficacy results\(^a\)

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>Ruxolitinib (n=110)</th>
<th>Best available therapy (n=112)</th>
<th>(P)-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary analysis (Week 32)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary composite endpoint(^*) (%)</td>
<td>20.9</td>
<td>0.9</td>
<td>(&lt;0.001)</td>
</tr>
<tr>
<td>Components of primary endpoint (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hematocrit control</td>
<td>60.0</td>
<td>19.6</td>
<td>NA</td>
</tr>
<tr>
<td>(\geq 35%) reduction in spleen volume</td>
<td>38.2</td>
<td>0.9</td>
<td>NA</td>
</tr>
<tr>
<td>Complete hematologic remission(^*) (%)</td>
<td>23.6</td>
<td>8.9</td>
<td>0.003</td>
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</tbody>
</table>

**Probability of maintaining response achieved**

**at primary analysis for \(\geq 80\) weeks**

| Components of primary endpoint (%)                                      |                      |                                |             |
|-------------------------------------------------------------------------|----------------------|                                |             |
| Hematocrit control                                                     | 89.0                 | NA                             | NA          |
| \(\geq 35\%\) reduction in spleen volume                               | NA\(^a\)             | NA                             | NA          |
| Complete hematologic remission\(^*\) (%)                                | 69.0                 | NA                             | NA          |

**Notes:** As reported by Vannucchi et al\(^10\) and Verstovsek et al.\(^10\) Defined as hematocrit control without phlebotomy eligibility (hematocrit \(>45\%\) and \(\geq 3\) percentage points higher than baseline or \(>48\%\), whichever was lower) and a \(\geq 35\%\) spleen volume reduction from baseline.\(^2\) Defined as hematocrit control without phlebotomy eligibility and platelet count \(\leq 400\times10^9/L\) and white blood cell count \(\leq 10\times10^9/L\).\(^3\) None of the patients in the ruxolitinib arm who achieved \(\geq 35\%\) reduction in spleen volume at Week 32 lost their response at the time of data cutoff for the 80-week analysis.

**Abbreviation:** NA, not available.
only one patient lost the response.109 The primary endpoint components were also durable; the probability of maintaining hematoctrit control without phlebotomy was 89%, and no patients who achieved ≥35% spleen volume reduction at Week 32 lost their response.109 Furthermore, patients who achieved a complete hematologic response in the primary analysis had a 69% probability of maintaining this response for ≥80 weeks after their initial response.109

Ruxolitinib treatment has also been associated with a decrease in JAK2V617F allele burden in patients with PV. In RESPONSE, the mean JAK2V617F allele burden changes from baseline in the ruxolitinib arm were −12.2% and −34.7% at Weeks 32 and 112, respectively.22 In contrast, patients randomized to receive the best available therapy had a mean 1.2% increase in JAK2V617F allele burden at Week 32.22 A single-center study of 22 patients with PV (n=11) or ET (n=11) reported mean changes in JAK2V617F allele burden of −19% and −28% after 36 months and 60 months of treatment with ruxolitinib, respectively.110

**Patient-reported outcomes**

Patient-reported outcomes favored ruxolitinib over the best available therapy in Phase III RESPONSE trial, with consistent improvements in the MPN-SAF, European Organisation for Research and Treatment of Cancer Quality-of-Life Questionnaire Core 30, Pruritus Symptom Impact Scale, and Patient Global Impression of Change instruments, compared with little improvement or worsening observed with the best available therapy (Table 2).22 Notably, even patients who did not achieve hematocrit control with ruxolitinib at Week 32 reported a higher degree of symptom improvement as evaluated by the MPN-SAF compared with patients who received the best available therapy (38% vs 4%, respectively).107

The randomized, multicenter, double-blind, double-dummy, Phase IIIb RELIEF trial was conducted in patients receiving a stable dose of hydroxyurea who were generally well controlled but reported disease-associated symptoms, comparing the change in PV-related symptom burden in patients continuing their hydroxyurea therapy with those switching to ruxolitinib treatment.111 There was a nonsignificant trend toward a greater improvement in the MPN-SAF cytokine cluster TSS (TSS-C) with ruxolitinib compared with hydroxyurea (proportion of patients achieving ≥50% reduction from baseline in TSS-C at Week 16: 43.4% vs 29.6%, respectively; primary endpoint).111 Additionally, there was a nonsignificant trend toward improvement in individual TSS-C symptoms with ruxolitinib compared with continued hydroxyurea in the RELIEF trial.111

### Table 2 Patient-reported outcomes in the RESPONSE triala

<table>
<thead>
<tr>
<th>Patient-reported outcome measure</th>
<th>Ruxolitinib</th>
<th>Best available therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with ≥50% reduction from baseline in MPN-SAF symptom severity scores at Week 32 (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MPN-SAF TSSb</td>
<td>49</td>
<td>5</td>
</tr>
<tr>
<td>Cytokine symptom clusterc</td>
<td>64</td>
<td>11</td>
</tr>
<tr>
<td>Hyperviscosity symptom clusterd</td>
<td>37</td>
<td>13</td>
</tr>
<tr>
<td>Splenomegaly symptom cluster*</td>
<td>62</td>
<td>17</td>
</tr>
<tr>
<td>Mean change from baseline in EORTC QLQ-C30 scores at Week 32*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Global health status/QoL</td>
<td>10.9</td>
<td>−4.8</td>
</tr>
<tr>
<td>Physical functioning</td>
<td>6.4</td>
<td>−1.5</td>
</tr>
<tr>
<td>Role functioning</td>
<td>5.3</td>
<td>−0.4</td>
</tr>
<tr>
<td>Emotional functioning</td>
<td>7.9</td>
<td>1.0</td>
</tr>
<tr>
<td>Cognitive functioning</td>
<td>4.2</td>
<td>−3.3</td>
</tr>
<tr>
<td>Social functioning</td>
<td>7.7</td>
<td>−0.4</td>
</tr>
<tr>
<td>Mean change from baseline on the Pruritus Symptom Impact Scale at Week 32*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>How severe was PV-related itching during the past 7 days</td>
<td>−2.2</td>
<td>0</td>
</tr>
<tr>
<td>How bothered by PV-related itching during the past 7 days</td>
<td>−2.0</td>
<td>0</td>
</tr>
<tr>
<td>How much PV-related itching interfered with daily life during the past 7 days</td>
<td>−1.5</td>
<td>0.3</td>
</tr>
<tr>
<td>How bothered by PV-related itching during the past 24 hours</td>
<td>−1.9</td>
<td>−0.1</td>
</tr>
<tr>
<td>How much PV-related itching interfered with daily life during the past 24 hours</td>
<td>−1.4</td>
<td>0.3</td>
</tr>
</tbody>
</table>

**Patient Global Impression of Change from baseline at Week 32 (%)**

| Very much improved | 32 | 4 |
| Much improved | 36 | 9 |
| Minimally improved | 11 | 21 |
| No change | 6 | 42 |
| Minimally worse | 1 | 13 |
| Much worse | 0 | 4 |
| Very much worse | 0 | 0 |

**Notes:** As reported by Vannuchi et al,22 all analyses included patients with assessments at baseline and Week 32. Sum of all 14 MPN-SAF symptom severity scores. Sum of cytokine-related MPN-SAF symptom severity scores (tiredness, itching, muscle aches, night sweats, and sweating while awake). Sum of hyperviscosity-related MPN-SAF symptom severity scores (headache, concentration problems, dizziness, skin redness, vision problems, ringing in ears, and numbness/shaking in hands/feet). Sum of splenomegaly-related MPN-SAF symptom severity scores (early satiety and abdominal discomfort). The EORTC QLQ-C30 global health status/QoL, and functional scales were linearly transformed to a 0–100 scale; higher scores represented a better level of functioning.111,112 Included patients with assessments at both baseline and Week 32. Patients responded to each question on a scale of 0 (not at all) to 10 (worst imaginable).

**Abbreviations:** EORTC QLQ-C30, European Organisation for Research and Treatment of Cancer Quality-of-life Questionnaire Core 30; MPN-SAF, Myeloproliferative Neoplasm Symptom Assessment Form; MPN-SAF TSS, MPN-SAF total symptom score; PV, polycythemia vera; QoL, quality-of-life.

### Safety and tolerability

The most common nonhematologic adverse events at Week 32 in patients who received ruxolitinib in the Phase III RESPONSE trial were headache, diarrhea, pruritus, and fatigue (Table 3).22,112 The most frequent grade 3 or 4 nonhematologic...
adverse events in the ruxolitinib arm were dyspnea, asthenia, abdominal pain, headache, muscle spasms, and pruritus. Long-term treatment data indicated that the incidences of the most common nonhematologic adverse events were similar at Week 48 and in the 80-week analysis (headache, 20.9% and 21.8%; diarrhea, 20.0% and 19.1%; pruritus, 20.0% and 17.3%; fatigue, 17.3% and 17.3%, respectively).112

Hematologic adverse events were primarily grade 1 or 2 at both the primary analysis (Table 3) and the 80-week analysis in the RESPONSE trial.12 The most common grade 3 or 4 hematologic adverse events in both randomized treatment arms were lymphopenia and thrombocytopenia. The US label for ruxolitinib indicates that thrombocytopenia, anemia, and neutropenia should be managed by dose reduction/interruption or transfusion.99

At Week 32, the rate of any grade 3 or 4 adverse events per 100 patient-years was lower in the ruxolitinib arm (28.8) than in the best available therapy arm (44.0) in the RESPONSE trial.22

Given the risks associated with thromboembolic events and disease transformation in patients with PV,14 it is important to consider such events when evaluating any new treatment option. The incidence of all-grade thromboembolic events at Week 32 was 0.9% in the ruxolitinib arm and 5.4% in the best available therapy arm (grade 3 or 4, 0.9% vs 1.8%, respectively).22 At the time of data cutoff in the primary analysis of RESPONSE, three patients in the ruxolitinib arm and one patient in the best available therapy arm developed MF after randomization; one patient in the ruxolitinib arm also developed AML. Two additional patients randomized to best available therapy experienced MF after cross over to ruxolitinib, one of whom progressed to AML.22

Other important adverse events of interest for patients being treated with ruxolitinib are nonmelanoma skin cancer and infections. At Week 32 in the RESPONSE trial, four patients in the ruxolitinib arm when compared with two patients in the best available therapy arm experienced newly diagnosed nonmelanoma skin cancer. All but one of these patients (best available therapy arm) had a history of precancerous skin lesions or nonmelanoma skin cancer.22 Melanoma skin cancer was diagnosed in zero patients in the ruxolitinib arm and one patient in the best available therapy arm during randomized treatment. The US prescribing information recommends periodic skin examinations for patients treated with ruxolitinib.99 In the RESPONSE trial, the incidence of any infection was similar in the ruxolitinib arm and the best available therapy arm at Week 32 (any grade, 41.8% vs 36.9%, respectively; grade 3 or 4, 3.6% vs 2.7%). However, herpes zoster infection was observed in 6.4% of patients in the ruxolitinib arm (all grade 1 or 2) at Week 32 compared to 0% of patients in the best available therapy arm.22 The US prescribing information states that patients should be advised about the early signs and symptoms of herpes zoster infection and to seek treatment as early as possible if herpes zoster infection is suspected.99

A long-term safety and tolerability evaluation of ruxolitinib was conducted in 241 patients (457 patient-years) who were resistant to or intolerant of hydroxyurea pooled from a Phase II study111 and RESPONSE (randomized and crossover patients).22,114 The most common nonhematologic adverse events were diarrhea, headache, and pruritus.114 Frequent grade 3 or 4 nonhematologic adverse events included dyspnea, herpes zoster, abdominal pain, back pain, headache, fatigue, and pyrexia.114 Hematologic adverse events were primarily grade 1 or 2; grade 3 or 4 anemia and thrombocytopenia were each reported in 3.7% of patients.114 Overall, seven patients had disease transformation to MF, and two patients had disease transformation to AML,114 a rate consistent with prior publications in similar patient populations with PV.21,84,115 The rate of thromboembolic events in the ruxolitinib group of this pooled analysis was 2.2/100 patient-years, whereas the rate in RESPONSE patients during randomized treatment with best available therapy was 8.2/100 patient-years.114

<table>
<thead>
<tr>
<th>Table 3 RESPONSE Phase III trial adverse eventsa</th>
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<tr>
<td>Adverse event (%)</td>
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<tr>
<td>All grade</td>
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<tr>
<td>Nonhematologicc</td>
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<tr>
<td>Headache</td>
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<tr>
<td>Diarrhea</td>
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<tr>
<td>Fatigue</td>
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<td>Pruritus</td>
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<tr>
<td>Dizziness</td>
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<tr>
<td>Muscle spasms</td>
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<tr>
<td>Dyspnea</td>
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<tr>
<td>Abdominal pain</td>
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<tr>
<td>Anemia</td>
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<tr>
<td>Hematologicc</td>
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<tr>
<td>Anemia</td>
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<tr>
<td>Thrombocytopenia</td>
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<tr>
<td>Lymphopenia</td>
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<tr>
<td>Leukopenia</td>
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<tr>
<td>Neutropenia</td>
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</tbody>
</table>

In Phase IIIb RELIEF trial, adverse events in the ruxolitinib arm were primarily grade 1 or 2. The most frequent nonhematologic adverse events were fatigue (20.4% [ruxolitinib arm] vs 10.7% [hydroxyurea arm]), headache (16.7% vs 5.4%), and dizziness (13.0% vs 8.9%). Two patients receiving ruxolitinib had grade 3 or 4 neutropenia.

Place in therapy and ongoing trials
Ruxolitinib is approved by the FDA for the treatment of patients with PV who have had an inadequate response to or are intolerant of hydroxyurea. Further research will be important for identifying other populations of patients with PV who may benefit from treatment with ruxolitinib. RESPONSE 2 (ClinicalTrials.gov identifier: NCT02038036) is an ongoing, randomized, open-label, Phase IIIb clinical trial designed to evaluate the efficacy and safety of ruxolitinib compared with the best available therapy in patients with PV without splenomegaly who are resistant to or intolerant of hydroxyurea and require phlebotomy. The primary endpoint is achievement of hematocrit control at Week 16 that is maintained through Week 28, together with no phlebotomy eligibility from Weeks 4 to 28. Patients who are randomized to the best available therapy and do not meet the primary endpoint are allowed to cross over to ruxolitinib at or after Week 28. Safety and durability of response will be evaluated through Week 52.

Conclusion
Ruxolitinib is the only approved treatment option designed to target the constitutively active JAK/STAT signaling pathway in patients with PV. Ruxolitinib improves hematocrit control without phlebotomy, improves blood cell counts, and reduces the enlarged spleen size. Accumulating evidence suggests that ruxolitinib may also ameliorate PV-related symptoms in patients who are resistant to and/or intolerant of hydroxyurea. Most adverse events were grade 1 or 2, and 82.7% of patients continued on treatment for ≥80 weeks in Phase III RESPONSE trial. Nonmelanoma skin cancer has been observed with ruxolitinib treatment, and periodic skin examinations should be performed. Herpes zoster infection rates (all grade 1 or 2) were higher with ruxolitinib compared with the best available therapy in the RESPONSE trial, and patients receiving ruxolitinib should be advised about the early signs and symptoms of infection and instructed to seek treatment as early as possible if suspected. Rates of disease transformation to MF and AML observed with ruxolitinib are similar to those previously published in similar patient populations with PV.

Collectively, clinical trial data indicate that ruxolitinib is an effective treatment option for many patients with PV who are resistant to and/or intolerant of hydroxyurea. Furthermore, the potential benefits of ruxolitinib are unique when compared with the traditional treatment options because ruxolitinib may alleviate the PV-related symptom burden and improve the QoL. Updated treatment guidelines will be important for educating physicians about using ruxolitinib for the treatment of patients with PV.

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Disclosure
KV is an employee of Incyte Corporation. SV participated in advisory boards and received research funding from Incyte Corporation. JJK served as a consultant for Incyte Corporation and Novartis and received a travel grant and research funding paid by Novartis to Hôpital Saint-Louis. The authors report no other conflicts of interest in this work.

References


