Retrospective Comparison of Patient Outcomes on Pacritinib Versus Ruxolitinib in Patients with Myelofibrosis and Thrombocytopenia

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BACKGROUND

- Pacritinib is a JAK2/IRAK1 inhibitor¹ approved by the Food and Drug Administration (FDA) in the United States for patients with myelofibrosis (MF) and thrombocytopenia.
- Unlike the JAK1/2 inhibitor ruxolitinib, which must be dose-reduced or held in patients with thrombocytopenia, pacritinib has been studied at full dose regardless of platelet count

OBJECTIVE

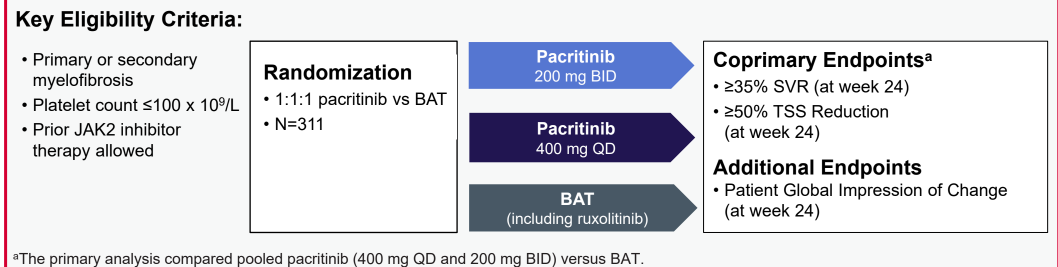
• To retrospectively analyze outcomes in patients treated with pacritinib versus ruxolitinib as part of the phase 3 PERSIST-2 study.

METHODS

Study Design

- In PERSIST-2, patients were randomized 1:1:1 to pacritinib 200 mg twice daily (BID), pacritinib 400 mg once daily (QD), or best available therapy (BAT)², as shown in Figure 1.
- 45% of patients on BAT received ruxolitinib.

Figure 1. PERSIST 2 Study Design



BAT=best available therapy; BID=twice daily; JAK2=Janus associated kinase 2; QD=once daily; SVR=spleen volume reduction; TSS=total symptom

- This analysis focuses on the approved dose of 200 mg BID dose for pacritinib and on patients who received ruxolitinib as BAT (BAT=RUX) prior to week 24.
- Safety analyses were based on all treated patients.
- Efficacy analyses were based on the intention-to-treat (ITT) population randomized at least 22 weeks prior to study end. The modified Total Symptom Score (mTSS) was used to assess MF symptoms.³
- Survival analysis was based on ITT for the pacritinib arm and treated patients for the ruxolitinib group.
- The Fisher's Exact test was used to describe response differences. Logistic or Cox proportional hazard regression models were used to adjust for baseline differences.

RESULTS

Patient Characteristics

- Safety analysis included 106 patients on pacritinib and 44 on ruxolitinib.
- ITT efficacy analysis (patients randomized ≥22 weeks prior to study end) included 74 on pacritinib and 32 on ruxolitinib.
- Baseline characteristics were similar between groups, including median platelet count $(55 \text{ vs } 61 \text{ x } 10^9\text{/L})$ and percentage receiving RBC transfusion (46% vs 43%).
- Patients in the ruxolitinib group were more likely to have prior JAK inhibitor exposure and PS ≥2.
- Presented at the Society of Hematologic Oncology September 28 October 1, 2022 Houston, TX

Table 1. Baseline Patient and Disease Characteristics

67 44 (42%) 12 (11%)	68 15 (34%) 10 (23%)
12 (11%)	
	10 (23%)
55	61
47 (44%)	17 (39%)
9.7	9.9
49 (46%)	19 (43%)
48 (45%)	27 (61%)
82 (77%)	22 (50%)
29 (27%)	12 (27%)
51 (48%)	32 (73%)
	47 (44%) 9.7 49 (46%) 48 (45%) 82 (77%) 29 (27%)

ion was not available for all patients in the safety population DIPSS=Dynamic International Prognostic Scoring System; EGOC=Eastern Cooperative Oncology Group; JAKi=Janus associated kinase inhibitor; PAC=pacritinib: PS=performance status: RBC=red blood cell: RUX=ruxolitinib

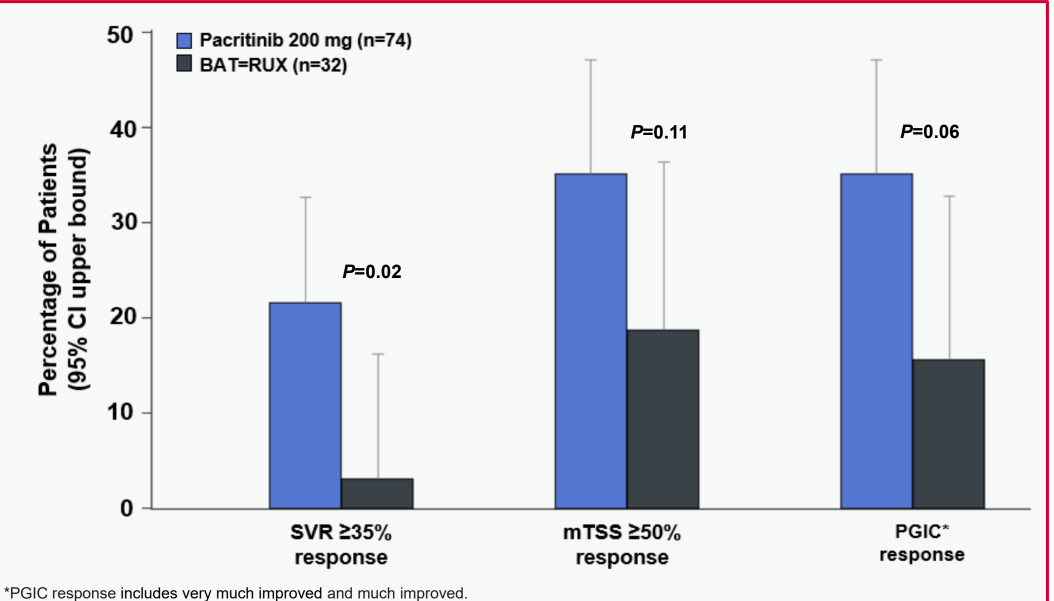
• The following differences in baseline characteristics were accounted for in the multiple regression model: percentage with grade 3 fibrosis, percentage with primary MF, percentage with ≥1% peripheral blasts, and percentage with prior JAK2 inhibitor use. • Median total daily dose of pacritinib was 400 mg [interquartile range (IQR): 400 – 400 mg] and ruxolitinib was 10 mg (IQR: 10 – 20 mg).

Efficacy

• Patients treated with pacritinib vs ruxolitinib achieved higher rates of spleen volume reduction (SVR) (22% vs 3%, P=0.02) and mTSS response (35% vs 19%; P=0.11) at week 24 (Figure 2)

• A greater percentage of patients on pacritinib reported "much" or "very much" improved symptoms (35% vs 16%, *P*=0.06); (**Figure 2**).

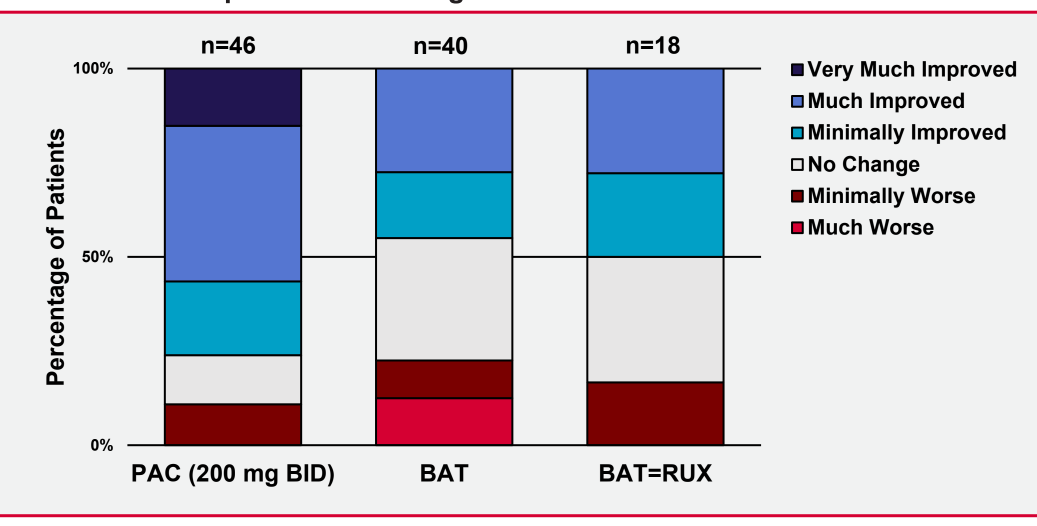
Figure 2. Efficacy Endpoints (Week 24)



PGIC=Patient Global Impression of Change; SVR=spleen volume response; TSS=total symptom score

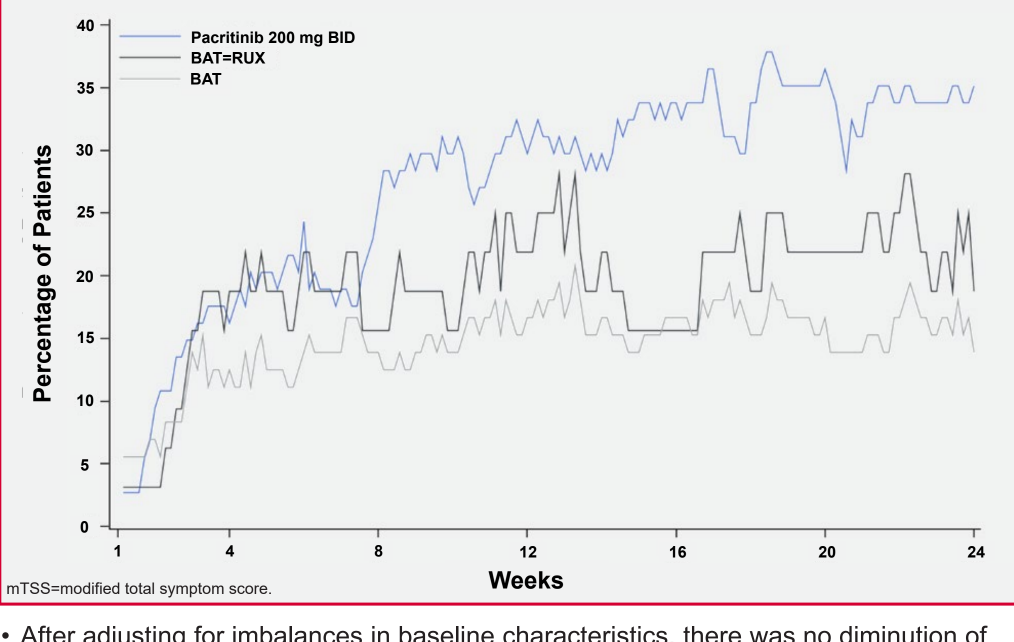
• Among ruxolitinib-treated patients with an available Patients' Global Impression of Change measure at week 24, 50% reported either no improvement or worsening symptoms, while 76% of pacritinib-treated patients reported improvement (Figure 3).

Figure 3. Patient-reported Change in MF Symptoms at Week 24 Based on **Patient Global Impression of Change**



• Rolling 7-day mTSS data shows rapid onset of symptom response by week 4, with ongoing improvement through weeks 12-24 (Figure 4).

Figure 4. Percentage of Patients with ≥50% Reduction in mTSS Score Based on **Rolling 7-day Assessment**

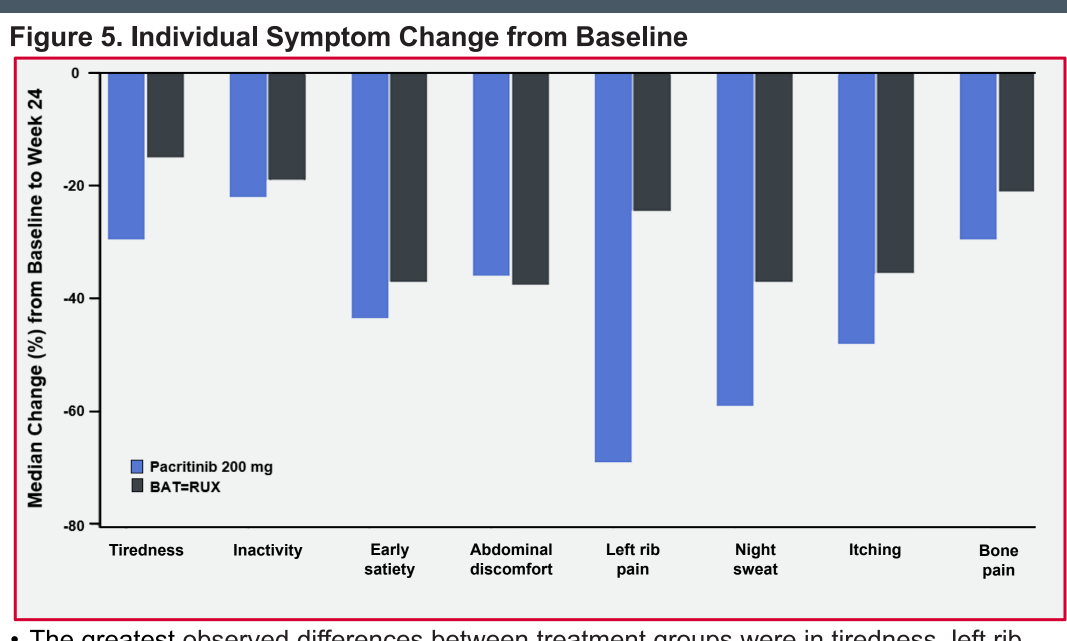


RESULTS

• After adjusting for imbalances in baseline characteristics, there was no diminution of treatment effect on SVR or mTSS.

• The hazard ratio for survival on pacritinib vs ruxolitinib was 0.71 (95% CI: 0.26-1.96). After adjusting for baseline imbalances between groups, the hazard ratio for survival on pacritinib vs ruxolitinib was 0.46 (95% CI: 0.15-1.43).

 Patients on the pacritinib arm had greater percentage reduction in most MF symptoms compared to patients who received ruxolitinib as BAT (Figure 5).



pain, night sweats, and itching.

Safety

- did bleeding events (Table 2).

Table 2. Adverse event (AE) Overview by Treatment Group

Patients with ≥1 AE, n (%)	PAC n=106	BAT=RUX n=44	
Any AE	100 (94%)	41 (93%)	
Fatal AE	8 (8%)	5 (11%)	
AE requiring dose reduction	13 (12%)	5 (11%)	
AE requiring drug withdrawal	16 (15%)	7 (16%)	
Withdrawal due to thrombocytopenia	2 (2%)	1 (2%)	
Withdrawal due to anemia	3 (3%)	0	
Hemorrhagic AE	45 (43%)	18 (41%)	
Cardiac AE	34 (32%)	10 (23%)	
Additional comparative safety data on pacritinib and ruxolitinib has bee	en previously presented. ⁴		
CONC	LUSIONS		

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REFERENCES: 1. Singer JW et al. J Exp Pharmacol. 2016;8:11-19. 2. Mascarenhas J, et al. JAMA Oncol. 2018;4(5):652-659. 3. Mesa RA et al. J Clin Oncol. 2013; 31(10):1285-1292. 4. Pemmaraju N et al. Risk-adjusted safety analysis of pacritinib in patients with myelofibrosis. Poster (7058) presented at ASCO Annual Meeting; June 2022, Chicago IL.



• The greatest observed differences between treatment groups were in tiredness, left rib

• Overall and fatal adverse events occurred at similar rates on pacritinib vs ruxolitinib, as

• Cardiac events occurred more commonly on pacritinib, though the difference was largely due to higher rates of grade 1 peripheral edema on pacritinib.

• There were low rates of herpes zoster reactivation (n=0 vs 1), fungal skin infection (n=0 vs 1), pulmonary aspergillosis (n=1 vs 0), deep venous thrombosis (n=0 vs 1), and pulmonary embolism (n=1 vs 0) on pacritinib and ruxolitinib, respectively.

• Pacritinib, administered at the full dose of 200 mg BID, yielded higher response rates and a similar safety profile compared to lower-dose ruxolitinib in patients with MF who have moderate or severe thrombocytopenia.

